Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	H H H	% ident	length (nt)
1 20	9	1 5089	4679	gi 511839	ORF1 [Staphylococcus bacteriophage phi 11]	100	100	411
1 149	m 	1 2032	1577	pir B49703 B497 int	lint gene activator RinA - bacteriophage phi 11	100	100	456
149	- - -	2109	1912	gi 166161 	Bacteriophage phi-11 int gene activator [Staphylococcus acteriophage phi	100	100	198
1 349	- 5	1 558	409	gi 166159	Integrase (int) [Staphylococcus bacteriophage phi 11]	100	100	150
398	1	1372	707	gi 166159	Integrase (int) [Staphylococcus bacteriophage phi 11]	100	66	999
398	7	1 783	1001	gi 455128	excisionase (xis) {Staphylococcus bacteriophage phi 11}	100	1000	219
502	4	1 1914	1744	gi 1204912	H. influenzae predicted coding region H10660 [Haemophilus influenzae]	100	71	171
849		7	262	gi 1373002	polyprotein [Bean common mosaic virus]	100	46	261
1349		772	140	gi 143359	protein synthesis initiation factor 2 (infB) [Bacillus subtilis] gi 49319 IF2 gene product [Bacillus subtilis]	100	82	138
1 2880		1 21	1 308	gi1862933	protein kinase C inhibitor-I [Homo sapiens]	100	86	288
3085	-	1 428	216	gi 1354211	PET112-like protein [Bacillus subtilis]	100	1000	213
4168	7	571	398	gi 1354211	PET112-like protein [Bacillus subtilis]	100	100	174
331	. –	7	247	gi 426473	nusG gene product [Staphylococcus carnosus]	1 86	95	246
1 207	7	1272	1463	gi 460259	enolase (Bacillus subtilis]	97	06	192
331	- 1	395	850	gi 581638	[L1] protein (Staphylococcus carnosus)	97	93	456
366		39	215	gi 166161 	Bacteriophage phi-11 int gene activator (Staphylococcus acteriophage phi	97	95	1771
1 680	3	718	936	gi 426473	InusG gene product [Staphylococcus carnosus]	97	97	219
3578		1 284	144	gi 1339950	large subunit of NADH-dependent glutamate synthase (Plectonema boryanum)	97	1 61	141
157		321	518	gi 1022726	unknown [Staphylococcus haemolyticus]	96	88	198
205	133	116470	116147	gi 1165302		1 96	91	324
3919		4 8	1 401	gi 871784	Clp-like ATP-dependent protease binding subunit [Bos taurus]	1 96	81	354
4133		830	417	gi 1022726	unknown [Staphylococcus haemolyticus]	1 96	84	414
4168		1 708	355	gi 1354211	PET112-like protein [Bacillus subtilis]	96	95	354
4207		312	157	gi 602031 		96	98	156
++	+-				<u>+</u>	+	+	+

Table

S. aureus - Putative coding regions of novel proteins similar to known proteins

4227 2 152 331 911871784 416 1 570 286 9411022728 22 1 858 430 94151070 82 6 8794 9114 piri/360008 154 9 9280 7838 9411354211 186 3 2798 2055 9411354211 205 7 5017 4793 941144245 205 7 5017 4793 94114445 205 7 5017 4793 941114245 205 7 5017 4793 941114245 205 7 5017 1997 9411177684 302 3 795 1097 9411177684 414 1 2 163 941171768 22 2 1028 723 941151106 23 5 5046 3310 941140516 60 4 815 1372 9411666116 205 18 10012 9536 941165386 414 1 3 737 955 941165386 414 1 1 1 1 205 18 10012 9536 9411667386 414 1 3 737 955 941167386 414 1 3 737 955 941167386 414 1 3 737 955 941167386 414 1 3 737 955 941167386 414 1 3 737 955 941167386 414 1 3 737 955 941167386 414 1 3 737 955 941167386	1.7600	Clp-like ATP-dependent protease binding subunit [Bos taurus]	96		
1 570 286 91 1 638 4036 91 6 8794 9114 91 9 9280 7838 91 1 5017 4793 91 1 11365 10991 91 1 579 1523 91 1 579 1523 91 2 125 127 91 3 737 955 91 4 815 1372 91 8 1378 2542 91 1 10012 9536 91 1 1 1378 191 1 1 1378 191 1 1 1 191 91 1 1 1 1 191 1 1 1 1 191 1 1 1 1 191 1 1 1 1 191 1 1 1 191 1 1 1 191 1 1 1 191 1 1 1 191 1 1 1 191 1 1 1 191 1 1 1 191 1 1 191				81	180
1 858 430 91 6 8794 9114 pi 9 9280 7838 91 3 2798 2055 91 5 4406 4014 91 7 5017 4793 91 1 579 1523 91 1 579 1523 91 1 579 1523 91 1 579 163 pi 2 125 277 91 2 126 723 91 3 5046 3310 91 4 815 1372 91 1 8 10012 9536 91 1 10012 9536 91 1 1 1 1 1 1 1 1 1		unknown (Staphylococcus haemolyticus)	96	84	285
7 4362 4036 91 6 8794 9114 91 9 9280 7838 91 5 4406 4014 91 7 5017 4793 91 1 579 1097 91 1 579 1523 91 1 579 1523 91 1 5 1028 723 91 2 1028 723 91 3 737 955 91 4 815 1372 91 4 815 1372 91 6 815 1372 91 737 955 91		UreG [Staphylococcus xylosus]	95	88	429
6 8794 9114 ppi 9 9280 7838 gi 1 2 4406 4014 gi 7 5017 4793 gi 1 11365 10991 gi 1 1579 1523 gi 1 579 1523 gi 1 5 1028 723 gi 2 125 277 gi 3 795 1372 gi 4 815 1372 gi 8 815 1372 gi 9 815 1372 gi 1 8 8 8 8 8 1 8 8 8 8 1 8 8 8 8 1 9 9 9 1 9 9 1 9	T — T -	urease gamma subunit (Staphylococcus xylosus)	95	79 1	327
9 9280 7838 91 3 2798 2055 91 7 5017 4793 91 21 11365 10991 91 3 7288 6644 8p 3 795 1097 91 1 579 1523 91 2 125 177 91 3 795 9536 91 4 815 1372 91 4 815 1372 91 6 815 1372 91 7 8 8 6 91 8 8 8 8 8 91 9 9 9 91	54211	ribosomal protein S7 - Bacillus stearothermophilus	95	83	321
3 2798 2055 91 5 4406 4014 91 7 5017 4793 91 3 795 1097 91 1 579 1523 91 1 2 164 191 1 2 164 191 1 2 164 191 1 2 164 191 2 125 277 91 3 5046 3310 91 4 815 1372 91 4 815 1372 91 7 3378 2542 91 8 737 955 91	117500	PET112-like protein (Bacillus subtilis)	95	92	1443
5 4406 4014 91 7 5017 4793 91 21 11365 110991 91 3 7288 6644 8p 1 579 1523 91 1 579 1523 91 2 125 127 91 5 5046 3310 91 6 815 1372 91 7 815 1372 91 8 8 8 8 8 91 9 9 9 91	11514656	serine O-acetyltransferase [Staphylococcus xylosus]	95	87	744
7 5017 4793 491	1142462	ribosomal protein S11 (Bacillus subtilis)	95	85	. 868
21	1142459	initiation factor 1 [Bacillus subtilis]	95	84	225
5 7288 6644 8p 3 795 1097 9i 1 579 1523 9i 1 2 163 pi 2 125 277 9i 2 1028 723 9i 4 815 1372 9i 4 815 1372 9i 6 3378 2542 9i 737 955 9i	11044974	ribosomal protein L14 (Bacillus subtilis)	95	93	375
3 795 1097	P47995 YSEA_	HYPOTHETICAL PROTEIN IN SECA 5'REGION (ORF1) (FRAGMENT).	95	85 1	645
1 579 1523 (gi 1 2 163 pi 2 125 277 gi 5 5046 3310 gi 4 815 1372 gi 4 3378 2542 gi 3 737 955 gi	40186	homologous to E.coli ribosomal protein L27 [Bacillus subtilis] il143592 L27 libosomal protein [Bacillus subtilis] ir[C21895]C21895 ribosomal protein L27 - Bacillus subtilis p P05657 RL27_BACSU 50S RIBOSOMAL PROTEIN L27 (BL30) (BL24). il40175 L24 gene prod	95	68	303
1 2 163 pi 2 125 277 gi 2 1028 723 gi 4 815 1372 gi 4 3378 2542 gi 3 737 955 gi	11177684	chorismate mutase [Staphylococcus xylosus]	95	92	945
2 125 277 91 2 1028 723 91 5 5046 3310 91 4 815 1372 91 18 110012 9536 91 4 3378 2542 91 3 737 955 91	r C48396 C483	ribosomal protein L34 - Bacillus stearothermophilus	95	06	162
2 1028 723 91 5 5046 3310 91 4 815 1332 91 18 10012 9536 91 4 3378 2542 91 3 737 955 91	1276841	glutamate synthase (GOGAT) [Porphyra purpurea]	95	98	153
5 5046 3310 91 4 815 1372 91 18 10012 9536 91 4 3378 2542 91 3 737 955 91	1511069	Uref [Staphylococcus xylosus]	94	91	306
4 815 1372 gi 18 10012 9536 gi 4 3378 2542 gi 3 737 955 gi	1410516	urease alpha subunit (Staphylococcus xylosus)	94 -	85 -	1737
18 10012 9536 91 4 3378 2542 91 	666116	glucose kinase (Staphylococcus xylosus)	9.6	87	558
4 3378 2542 91	11044978	ribosomal protein S8 (Bacillus subtilis)	94	78	477
3 737 955 gi	557492	dihydroxynapthoic acid (DHNA) synthetase (Bacillus subtilis) gil143186 dihydroxynapthoic acid (DHNA) synthetase (Bacillus ubtilis)	46	85	837
	1467386	thiophen and furan oxidation [Bacillus subtilis]	94	7.7	219
426 3 2260 1823 gi 126:	11263908	putative (Staphylococcus epidermidis)	94	87	438
534 1 2 355 91 633	1633650	enzyme II(mannitol) [Staphylococcus carnosus]	94	84	354
1017 1 2 229 1911149	1149435	putative (Lactococcus lactis)	94	73	228
3098 1 330 184	1413952	ipa-28d gene product [Bacillus subtilis]	94	50 -	147

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

1	1		1			+	+	+11111111
Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
3232	-	630	316	gi 1022725	unknown [Staphylococcus haemolyticus]	94	84	315
42	S	5089	1 2259	pir B48396 B483	ribosomal protein L33 - Bacillus stearothermophilus	93	81	171
101	2	1745	1383	gi 155345	arsenic efflux pump protein [Plasmid pSX267]	93	82	363
205	124	12227	111865	sp P14577 RL16_	SOS RIBOSOMAL PROTEIN L16.	93	83	363
259	4	8291	1 5673	gi 499335	secA protein [Staphylococcus carnosus]	93	85	2619
275		2226	1114		enzyme II(mannitol) [Staphylococcus carnosus]	93	9.8	1113
444	9 -	1 6207	1 5773	gi 1022726	unknown [Staphylococcus haemolyticus]	93	81	435
491	-	152	622	gi 46912	ribosomal protein L13 (Staphylococcus carnosus)	93	88	471
1 607	9	1674	1 2033	gi 1022726	unknown [Staphylococcus haemolyticus]	93	83	360
653		973	488	gi 580890		93	77	486
1864	-	e -	1 194	gi 306553	ribosmal protein small subunit [Homo sapiens]	93	93	192
1 2997		1 28	300	gi 143390	carbamy1 phosphate synthetase [Bacillus subtilis]	93	82	273
3232	1 2	1 907	1 596	gi 1022725	unknown [Staphylococcus haemolyticus]	93	84	312
3761	7	1 794	621	19111022725	unknown [Staphylococcus haemolyticus]	93	88	174
16		. m	374	gi 142781 	putative cytoplasmic protein; putative (Bacillus subtilis) sp P37954 UVRB_BACSU EXCINUCLEASE ABC SUBUNIT B (DINA PROTEIN) FRAGMENT).	92	83	372
31		5915	6124	gi 1136430	KIAA0185 protein [Homo sapiens]	92	46	210
99	119	126483	127391	gi 467401	unknown [Bacillus subtilis]	95	80	606
69	9	5882	6130	gi 530200	trophoblastin [Ovis aries]	95	53	249
145	m -	1 2568	1 2038	gi 1022725	unknown [Staphylococcus haemolyticus]	92	08	531
171	m	1 2760	1 2362	gi 517475	D-amino acid transaminase (Staphylococcus haemolyticus)	92	98	399
205	112	7495	1 6962	gi 49189	secY gene product [Staphylococcus carnosus]	92	85	534
1 205	119	110812	110255	gi 1044976		92	82	558
219	1	710	1 357	gi 1303812		95	88	354
344	m 	1575	1 1805	gi 1405474	CspC protein [Bacillus cereus]	95	85	231
669	-	50	361			95	81	342
1343	- 1	- 5	160	pir A45434 A454	r A45434 A454 ribosomal protein L19 - Bacillus stearothermophilus	92	84	159
	+							

Table

| % sim | % ident | length (nt) hsdM gene of EcoprrI gene product [Escherichia coli] pir18384371838437 hsdM | protein - Escherichia coli pir18096291809629 hypothetical protein A - | Escherichia coli (SUB 40-520) fructose-bisphosphate aldolase [Staphylococcus carnosus] pir|A49943|A49943 |large subunit of NADH-dependent glutamate synthase [Plectonema boryanum] [large subunit of NADH-dependent glutamate synthase [Plectonema boryanum] fructose-bisphosphate aldolase (EC 4.1.2.13) - taphylococcus carnosus adenylosuccinate lyase (PUR-B) [Bacillus subtilis] pir|C29326|WZBSDS orf2 downstream of glucose kinase [Staphylococcus xylosus] adenylosuccinate lyase (EC 4.3.2.2) - Bacillus ubtilis ATP synthase alpha subunit (Bacillus megaterium) ATP synthase subunit beta (Bacillus subtilis) |glcA gene product [Staphylococcus carnosus] |arsenic efflux pump protein [Plasmid pI258] |sucrose repressor [Staphylococcus xylosus] (S17 protein (AA 1-87) [Bacillus subtilis] ccpA gene product [Staphylococcus xylosus] hypothetical protein [Synechocystis sp.] |unknown [Staphylococcus haemolyticus] |unknown [Staphylococcus haemolyticus] |unknown [Staphylococcus haemolyticus] |unknown [Staphylococcus haemolyticus] |IMP dehydrogenase [Bacillus subtilis] |GMP synthetase (Bacillus subtilis) |CTP synthetase [Bacillus subtilis] [Eliscr [Staphylococcus xylosus] [unknown [Bacillus subtilis] [CodW [Bacillus subtilis] (strain TM300) match gene acession gi|1339950 |gi|1339950 gi|1022726 gi | 1022725 |gi|1022726 gi|1022725 |gi|1001376 1gi11072418 |gi|1226043 1gi11177685 |gi|407908 |gi|450688 1gi 1535349 |gi|142559 |gi|143012 1gi1467399 |gi|949974 |gi|143366 gi | 297874 |gi|385178 |gi|433991 |gi|143597 |gi|40149 Stop (nt) 1 2326 1 3826 1 3207 1 3150 1 4531 Start 1 2974 (nt) 1 8184 1 1606 1 1 21 1 412 ---Contig 10RF -Ω -S <u>۳</u> 'n <u>~</u> m ~ C

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

	111111		1111111	TOTAL CONTRACTOR OF THE PARTY.		-		1 1 1 1 1 1 1	
Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	. 	sim	% ident	length (nt)
1 1327		339	530	gi 496558	orfX [Bacillus subtilis]		91	1.1	192
2515	-	466	275	gi 511070	UreG (Staphylococcus xylosus)		91	85	192
1 2594		2	202	gi 146824	 beta-cystathionase [Escherichia coli]		91	75	201
3764	-	847	425	gi 1022725	unknown [Staphylococcus haemolyticus]		91	78	423
4011	-	127	495	gi 1022726	unknown [Staphylococcus haemolyticus]		91	97	369
4227		1	177	gi 296464	ATPase [Lactococcus lactis]		91	99	1771
42	8	815	1033	lgi1520401	catalase [Haemophilus influenzae]	-	06	98	219
51	8 -	3717	4607	1911580899	OppF gene product [Bacillus subtilis]		06	74	891
129	3	5317	4001	gi 1146206	glutamate dehydrogenase [Bacillus subtilis]		06	9/	1317
164	117	116628	16933	sp P05766 RS15_	30s Ribosomal protein s15 (BS18).		1 06	74	306
171	2	2983	2819	gi 517475	D-amino acid transaminase [Staphylococcus haemolyticus]		1 06	78	165
1 205	4	4497	3550	gi 142463	RNA polymerase alpha-core-subunit [Bacillus subtilis]		1 06	97	948
205	9	4748	4410	gi 1044989	ribosomal protein S13 [Bacillus subtilis]	-	06	73	339
1 205	110	7165	6404	gi 49189	secY gene product [Staphylococcus carnosus]		06	81	762
205	111	6645	6472	gi 49189	secY gene product [Staphylococcus carnosus]		06	78	174
205	127	13692	113345	gi 786157	Ribosomal Protein S19 [Bacillus subtilis]		1 06	62	348
205	,	115858	115496	gi 1165303	[L3 (Bacillus subtilis]		1 06	67	363
260	2	1 7023	5773	gi 1161380		-	1 06	78	1251
299	9	3378	3947	gi 467440 	phosphoribosylpyrophosphate synthetase [Bacillus subtilis] gi 40218 PRPP synthetase (AA 1-317) [Bacillus subtilis]		06	78	570
320	7	1025	1717	gi 312443	carbamoyl-phosphate synthase (glutamine-hydrolysing) [Bacillus aldolyticus]		06	75	693
330	4	1581	1769	gi 986963	beta-tubulin [Sporidiobolus pararoseus]		1 06	80	189
369		954	523	pir 834762 8347	L-serine dehydratase beta chain - Clostridium sp.		06	77	432
557		3	188	gi 1511589	M. jannaschii predicted coding region MJ1624 [Methanococcus jannaschii]		1 06	54	186
663	7	199	1200	gi 143786 	tryptophany1-tRNA synthetase (EC 6.1.1.2) [Bacillus subtilis] pir JT0481 YWBS tryptophantRNA ligase (EC 6.1.1.2) - Bacillus ubtilis		06	73	534
717		-	261	gi 143065	hubst [Bacillus stearothermophilus]	_	- 06	19	261
745	4	1059	865	gi 1205433	H. influenzae predicted coding region HI1190 (Haemophilus influenzae)		1 06	81	195
	-						-		

| % ident | length (nt) 11 83 9 / 22 91 75 17 84 9 / 80 80 74 80 82 20 80 74 75 80 70 68 83 89 68 83 # Sim 90 90 90 90 90 90 90 90 68 83 89 68 68 89 83 89 89 83 83 [phosphoglucose isomerase A (AA 1-449) [Bacillus stearothermophilus] ir[S15936]NUBSSA glucose-6-phosphate isomerase (EC 5.3.1.9) A - cillus |pir|(41902|(419 |arsenate reductase (EC 1.....) - Staphylococcus xylosus plasmid pSX267 | ir|S15936|NUBSSA glucose-6-phosphate isomerase (EC 5.3.1.9) A - cillus (formyltetrahydrofolate synthetase (FTHFS) (ttg start codon) (EC .3.4.3) ladenylosuccinate lyase (FUR-B) [Bacillus subtilis] pirlC29326|WZBSDS | adenylosuccinate lyase (EC 4.3.2.2) - Bacillus ubtilis phosphoglucose isomerase A (AA 1-449) [Bacillus stearothermophilus] S. aureus - Putative coding regions of novel proteins similar to known proteins [ATP-dependent protease binding subunit [Haemophilus influenzae] |Clp-like ATP-dependent protease binding subunit [Bos taurus] |ORF for L15 ribosomal protein [Bacillus subtilis] [glutamate racemase [Staphylococcus haemolyticus] triose phosphate isomerase [Bacillus subtilis] [IF2 (aa 1-741) [Bacillus stearothermophilus] [thermonuclease [Staphylococcus intermedius] ATP-binding protein [Streptococcus mutans] [cell division protein [Bacillus subtilis] |protein-dependent (Bacillus subtilis) [protease [Staphylococcus epidermidis] |ORF f729 [Escherichia coli] |unknown [Bacillus subtilis] [unknown [Bacillus subtilis] [Moorella thermoacetica] [YgeV [Bacillus subtilis] (ClpP [Bacillus subtilis] |YqeQ [Bacillus subtilis] [L3 [Bacillus subtilis] [L3 [Bacillus subtilis] | stearothermophilus stearothermophilus | match gene name acession |gi|1033122 |gi|1477776 |gi|1303812 |gi|1165303 1gi | 1377831 |gi|1070014 |gi|1165303 |gi|1205108 |gi|1303804 match |gi|143366 1gi1144816 1gi1871784 lgi | 153741 |gi|467458 |gi|216338 |gi|520574 |g1|396259 1gi1467327 1gi1460257 1gi147146 gil40046 |gi|39954 gil40046 132 | 116152 | 115823 1 1 | 1028 | 1750 14 | 3545 | 3757 7405 1 2407 | 2207 1 1 | 2124 | 1210 112755 (nt) 1 707 1 205 1 269 1 233 839 444 565 1 505 672 1 331 166 398 341 813 1 367 Contig |ORF | Start | ID |ID | (nt) | 113 | 7875 111667 1156 (1) 117 1 157 1 1 18 1 1 386 1 1 1 120 1054 | 1 | 579 1180 | 1 | 408 1 1 327 1111 | 1 | 352 | 1 | 11 11 + 3 4594 | 1 | 3 11 1 2 -3 ლ — 7 -112 1007 2951 3140 615 118 128 205 494 81 112 164 205 270 395

279 213 1089 471 330 201 516 837

504

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249 591 204

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162 231 723

267

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

1	Contig	ORF	Start	Stop (nt)	match acession	match gene name	# sim	% ident	length (nt)
1 100 214 GillSGG12 MATP synthase abbuilt games [Becillus aubtilla] 69 75 75 75 75 75 75 75 7	1 3020	1	06	362		protein (Bacillus	68	99	273
1 195 314 9 1180632 Mary synthese subbnilt geame [Bacillus subtilis] 89 182	3565		2	400		[Bacillus	68	75	399
1 2 400 4011000366 Respiratory nitrate reducence (Bacillus subtilis) 89 75 75 78 78 78 78 78 78	3586		105	314	gi 580832	ATP synthase subunit gamma [Bacillus subtilis]	68	82	210
1 2 4 600 4	1 3629	7 -	1 794	399		Respiratory nitrate reductase (Bacillus subtilis)	68	78	396
1 724 339 pill339960 large subunit of MADH-dependent glutamate synthase [Plectonoma boryanum] 69 156 156 151 151 152	3688		2	400	gi 1146206	glutamate dehydrogenase [Bacillus subtilis]	68	7.5	399
6 11 (428) 216 (911009366) Respiratory nitrate reductase [Bacillus subtilis] 69 71 7 11 (471) 301 (91149426 Puntative [Lactococcus haemolyticus] 69 76 8 11 (501) 302 (911022732) Unitroom (Staphylococcus haemolyticus) 69 76 1 (501) 302 (911022732) Unitroom (Staphylococcus haemolyticus) 69 76 1 (501) 267 (911022732) Unitroom (Staphylococcus haemolyticus) 69 76 1 (501) 302 (911022732) Unitroom (Staphylococcus wylosus) 68 76 1 (601) 302 (91104477) Indoordate (Staphylococcus wylosus) 68 76 1 (602) 4602 4722 (91104477) Indoordate (Staphylococcus wylosus) 68 77 2 (602) 472 (9110477) Indoordate (Staphylococcus wylosus) 68 77 2 (702) 472 (9110477) Indoordate (Staphylococcus wylosus) 68 77 2 (702) 472 (9110477) Indoordate (Staphylococcus wylosus) 68 77 3 (702) 472 (9110477) Indoordate (Staphylococcus wylosus) 68 77 4 (702) 472 (91104774) Indoordate (Staphylococcus wylosus) 68 77 <td>3699</td> <td></td> <td>1 794</td> <td>399</td> <td> gi 1339950</td> <td>large subunit of NADH-dependent glutamate synthase [Plectonema boryanum]</td> <td>68</td> <td>7.5</td> <td>1 968</td>	3699		1 794	399	gi 1339950	large subunit of NADH-dependent glutamate synthase [Plectonema boryanum]	68	7.5	1 968
7 11 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1) 71 (0.1)<	4016		1 428	216	gi 1009366	Respiratory nitrate reductase [Bacillus subtilis]	68	7.1	213
1 601 1002 49110027255 Unknown [Staphylococcus haemolyticus] 691 73 73 74 75 75 75 75 75 75 75	4177		471	301	gi 149426	[Lactococcus	68	9.	171
1 1320 162 1911022725 Unriknoom [Stephy]cooccus haemolyticus] 192 1320 1826 1911022725 Unriknoom [Stephy]cooccus xylosus] 198 176 198 176 198 177 177	4436		601	302	gi 1022725		68	80	300
1 188 848 848 849 849 849 849 840	4635		320	162	gi 1022725	unknown [Staphylococcus haemolyticus]	68	73	159
2 466 848 SpiP4222IICATA_CATALARE (EC I.11.1.6). 88 76 76 76 78 78 78 78	2	1 2	1330	2676	gi 520754	putative (Bacillus subtilis)	88	92	1347
5 6389 4722 491147177 alpha-D-1.44glucosidase [Staphylococcus xylosus] 88 77 88 77 88 18 1	1 42	1 2	1 468	848		CATALASE (EC 1.11.1.6).	88	92	381
15 1807 1816 1867 gji 1467411	1 53	2	6389	4722		alpha-D-1,4-glucosidase [Staphylococcus xylosus]	88	80	1668
3 376 943 qi 666116 qlucose kinase. Staphylococcus xylosus		!	i	118617		recombination protein [Bacillus subtilis]	88	77	009
2 1583 1245 gil44095 Teplication initiator protein [Listeria monocytogenes] 88 74 19 11514 11519 pir/A60663 A606 translation elongation factor Tu - Bacillus subtilis 88 77 82 1479 4391 gil167181 Serine/threonine kinase receptor [Brassica napus] 88 77 1 1 1569 8211 gil1022726 Unknown [Staphylococcus haemolyticus] 88 77 1 1 1 1 1 1 1 1	09	3	376	843	gi 666116	kinase (Staphylococcus	88	77	468
8 11514 12719 pir/A606631A606 translation elongation factor Tu - Bacillus subtilis 88 79 77 71 71 71 71 71 71	100	5	1583	1245	gi 44095	replication initiator protein [Listeria monocytogenes]	88	74	339
7 4179 4391	82			112719		elongation factor Tu - Bacillus	88	79	1206
8 7732 8232 gill022726 lunknown [Staphylococcus haemolyticus] 88 72 72 73 74 75 74 75 75 75 75 75	103	7	4179	4391		serine/threonine kinase receptor (Brassica napus)	88	77	213
2 308 2011 gill303804 Yqqo [Bacillus subtilis] 88 77 136 gill405446 transketolase [Bacillus subtilis] 88 72 136 gill118002 dihydropteroate synthase [Staphylococcus haemolyticus] 88 80 78 1428 12231 gill40053 phenylalanyl-tRNA synthetase alpha subunit [Bacillus subtilis] Bacillus subtilis 88 78 78 78 78 78 78 7	114	80	1732	8232	gi 1022726	unknown [Staphylococcus haemolyticus]	88	72	501
3 657 1136 gi 1405446 transketolase [Bacillus subtilis] 88 72 7 5871 6116 gi 1118002 dihydropteroate synthase [Staphylococcus haemolyticus] 88 78 78 78 78 78 78 7	118	7	308	2011		YqeQ [Bacillus subtilis]	88	77	1704
7 5871 6116 gill128002 dihydropteroate synthase [Staphylococcus haemolyticus]	141	m	1 657	1136	gi 1405446	transketolase [Bacillus subtilis]	88	72	480
3 1428 2231 gi 40053 phenylalanyl-tRNA synthetase alpha subunit (Bacillus subtilis) 88 80	148	7	5871	6116	gi 1118002	synthase	88	78	246
128 115027 114185 gi 1165306 LZ [Bacillus subtilis] 88 82	165	m 	1428	2231	gi 40053 	synthetase alpha subunit [Bacillus subtilis] henylalaninetRNA ligase (EC 6.1.1.20) alpha ain	88	08	804
1 1569 898		1	!	114185			88	82	843
1 2 1975 gi 452309 valy1-tRNA synthetase [Bacillus subtilis]	1 225		1569	868	gi 1303840	YqfS (Bacillus subtilis)	88	78	672
	235	-	2		gi 452309	valy1-tRNA synthetase (Bacillus subtilis)	88	9.2	1974

Table 2

495 1398 417 465 186 516 264 174 306 165 138 150 396 1431 300 231 351 408 339 366 336 | % sim | % ident | length (nt) 55 71 70 99 73 73 78 70 78 82 73 73 10 67 88 11 84 84 82 82 8.7 87 88 88 88 88 88 88 88 88 88 88 88 88 88 88 88 88 88 88 88 mannitol-specific enzyme-III (Staphylococcus carnosus) pirlJQ0088|JQ0088 pyruvate decarboxylase (E-1) beta subunit (Bacillus subtilis) gi/1377836 |valy1-tRNA synthetase [Bacillus stearothermophilus] sp|P11931|SYV_BACST phosphotransferase system enzyme II (EC .7.1.69), mannitol-specific, factor III - Staphylococcus carnosus splP178761PTMA_STACA_PTS_SYSTEM, MANNITOL-SPECIFIC IIA COMPONENT EIIA-MTL) (|pir|B47154|B471 |signal recognition particle 54K chain homolog Ffh - Bacillus subtilis pir184715418471 | signal recognition particle 54K chain homolog Ffh - Bacillus subtilis glutamate synthase (ferredoxin) [Synechocystis sp.] pir|S46957|S46957 |glutamate synthase (ferredoxin) [Synechocystis sp.] pir|S46957|S46957 | ir|S05348|R5BS20 ribosomal protein L20 - Bacillus earothermophilus VALYL-TRNA SYNTHETASE (EC 6.1.1.9) VALINE--TRNA LIGASE) (VALRS). glutamate synthase (ferredoxin) (EC 1.4.7.1) - ynechocystis sp. glutamate synthase (ferredoxin) (EC 1.4.7.1) - ynechocystis sp. |ribosomal protein L20 (AA 1-119) [Bacillus stearothermophilus] pyruvate decarboxylase E-1 beta subunit (Bacillus ubtilis) [pyrimidine nucleoside phosphorylase [Bacillus subtilis] |dihydropteroate synthase [Staphylococcus haemolyticus] |O-acetylserine(thiol) lyase (Spinacia oleracea) |dihydroxyacetone kinase [Citrobacter freundii] [nitrite reductase (nirB) [Bacillus subtilis] |glutamate dehydrogenase [Bacillus subtilis] [phosphomannomutase [Mycoplasma pirum] |unknown (Staphylococcus haemolyticus) |function umknown [Bacillus subtilis] (aconitase [Bacillus subtilis] ORF f746 [Escherichia coli] [CodX [Bacillus subtilis] [CodX [Bacillus subtilis] | match gene name acession 1gi11022726 |gi|1146206 | 8 | 6781 | 6482 |gi|1064791 1gi | 1118002 gi!1405454 |gi|558559 match |gi|143797 | 2968 |gi|153049 |gi|710018 |gi|401786 |gi|493083 1911606055 |gi|143378 gi | 515938 | 5 | 4570 | 6000 |gi|535350 gi | 535350 |gi|218277 gi | 515938 gil39963 13 | 2060 | 1566 1309 (nt) 1 324 214 399 1 419 399 1400 399 169 138 152 1 369 336 533 410 342 Contig | ORF | Start | ID | ID | (nt) | 1 3 | 2504 1 1824 4325 | 2 | 584 183 1 1 1 794 1 1 587 1341 | 2 | 170 1 1386 | 1 | 41 817 980 1 1 1 734 1 1 704 m -- ر د 1 - 3 --- 5 7 - 4 _ 1299 3089 2949 2984 3035 3917 4201 1386 4199 4274 4308 1000 339 443 25

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

3 1 1 2 1 1 2 1 1 1 1	4 2480				+	
11		0 gi 142992 	glycerol kinase (glpK) (EC 2.7.1.30) [Bacillus subtilis] pir B45866 B45866 glycerol kinase (EC 2.7.1.30) - Bacillus subtilis sp[P18157]GLPK_BACSU GLYCEROK KINASE (EC 2.7.1.30) (ATP:GLYCEROL - PHOSPHOTRANSFERASE) (GLYCEROKINASE) (GK).	68	72	897
1 2 2 1 2 2 1 1 2 5 6 4 4 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	3 9100	0 gi 467433	unknown [Bacillus subtilis]	87	62	288
2 1 1 2 2 1 1 2 2 1 1 1 2 2 1 1 1 2 2 1 1 1 2 2 1 1 1 2 2 1 1 1 2 2 1 1 1 1 2 2 1	5 2988	8 gi 556886	serine hydroxymethyltransferase [Bacillus subtilis] pir S49363 S49363 serine hydroxymethyltransferase - Bacillus ubtilis	87	1	1278
113 5 1 1 5 1 1 1 2 1 1 1 1 1 1 1 1 1 1 1 1	7 4032	gi 5	Unknown [Bacillus subtilis]	87	1 99	426
1 2 2 1 1 2 2 1 3 3 1 3 1 3 1 3 1 3 1 3	1 4559	9 gi 467460	unknown [Bacillus subtilis]	87	1 02	819
1 2 5 3 5 5 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	0 13810	0 gi 39954	IF2 (aa 1-741) [Bacillus stearothermophilus]	87	72	1101
1 23 23 - 1	4 2126	19i14	unknown [Bacillus subtilis]	87	1 8/	1023
125 23 1 1 1 1 1 1 1 1 1	2 1158	8 gi 143527	liron-sulfur protein [Bacillus subtilis]	87	177	825
125	7 2933	į	pir A27763 A277 succinate dehydrogenase (EC 1.3.99.1) flavoprotein - Bacillus subtilis	87	1 08 1	1785
125	2 11543	3 gi 1044972	ribosomal protein L29 [Bacillus subtilis]	87	1 82	240
	5 12607	7 gi 1165309	Sacillus subtilis]	87	75	699
	3 1107	7 gi 1177249	rec233 gene product [Bacillus subtilis]	87	104	927
236 3 1635	5 1333	3 gi 1146198	ferredoxin [Bacillus subtilis]	87	1 08 1	303
246 5 2585	5 2292	2 gi 467373	ribosomal protein S18 [Bacillus subtilis]	87	1 11	294
260 2 4189	9 3422	2 91 1161382		87	72	168
320 3 1696	6 2391	1 (gi1312443	carbamoyl-phosphate synthase (glutamine-hydrolysing) [Bacillus aldolyticus]	87	80	969
380 4 1165	5 1383	3 gi 142570	ATP synthase c subunit [Bacillus firmus]	87	1 08	219
414 4 900	1 1073	3 gi 467386	thiophen and furan oxidation [Bacillus subtilis]	8.1	77	174
425 2 1003	3 794	gi 1046166		87	1 69	210
448 1 1255	5 722	gi 405134	acetate kinase [Bacillus subtilis]	87	75	534
480 1 1	117	gi 142559	ATP synthase alpha subunit (Bacillus megaterium)	87	1 64 1	711
481 1 2	1 352	sp Q06797 RL1_B	50S RIBOSOMAL PROTEIN L1 (BL1).	87	12	351
7 2	1 955	gi 460911		87	1 28 1	597
677 3 934	1284	4 gi 460911	fructose-bisphosphate aldolase [Bacillus subtilis]	87	1 8 1	351
876 1 3	1 452	gi 1146247	asparaginyl-tRNA synthetase [Bacillus subtilis]	1 87	1 67 1	450

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop	match acession	match gene name	# sim	m % ident	l length (nt)
1376	-	426	1 214	gi 1065555	F46H6.4 gene product [Caenorhabditis elegans]	1 87	7 1 75	213
2206		е -	374	gi 215098	excisionase [Bacteriophage 154a]	87	7 1 72	372
2938	-		1 290	gi 508979	GTP-binding protein (Bacillus subtilis)	1 87	69 6	288
3081	1 2	126	308	gi 467399	IMP dehydrogenase [Bacillus subtilis]	1 87	7 1 72	183
3535	-		1 401	gi 1405454	aconitase [Bacillus subtilis]	1 87	08 1 4	399
4238		1 547	1 275	gi 603769	HutU protein, urocanase [Bacillus subtilis]	8	87 73	273
4	8	110427	8736	gi 603769	HutU protein, urocanase (Bacillus subtilis]	8	86 72	1692
22	9	4190	3738	 gi 410515	urease beta subunit [Staphylococcus xylosus]	8	86 73	453
54	- 5	1 2480	1 1572	 gi 289287	[UDP-glucose pyrophosphorylase [Bacillus subtilis]	8	02 98	606
124	m 	1 2336	1713	gi 556887	uracil phosphoribosyltransferase [Bacillus subtilis] pir S49364 S49364 uracil phosphoribosyltransferase - Bacillus ubtilis	60 1 1 1 1	86 74	624
148	8	1349	3448	gi 467458		8	86 75	1 2100
148	4	1 3638	3859	gi 467460	unknown [Bacillus subtilis]	80	86 1 73	1 222
152	e -	1340	1 2086	gi 1377835	Ipyruvate decarboxylase E-1 alpha subunit [Bacillus subtilis]	8	86 75	747
164	118	117347	119467	gi 1184680	polynucleotide phosphorylase [Bacillus subtilis]	8	86 72	2121
180	1 2	1 554	1159	gi 143467	ribosomal protein S4 (Bacillus subtilis]	8	86 1 80	909
205		1 2966	2592	gi 142464	ribosomal protein L17 [Bacillus subtilis]	80	77 98	375
205	126	13364	112990	gi 40107 	ribosomal protein L22 [Bacillus stearothermophilus] ir S10612 S10612 ribosomal protein L22 - Bacillus earothermophilus	. – –	86 75	375
246	7	3463	3140	gi 467375	ribosomal protein S6 [Bacillus subtilis]	80	02 98	324
299	<u>ه</u>	1196	1540	gi 39656	spoVG gene product [Bacillus megaterium]	8	02 198	345
299		3884	4345	gi 467440 	'phosphoribosylpyrophosphate synthetase [Bacillus subtilis] gi 40218 PRPP synthetase (AA 1-317) [Bacillus subtilis]	ω	86 78	1 462
304	5	1 2170	1 2523	gi 666983	[putative ATP binding subunit [Bacillus subtilis]	8	86 65	354
310	1 2	1487	1678	gi 1177684	chorismate mutase [Staphylococcus xylosus]	8	86 71	192
337	- 5	1 2086	3405	gi 487434	[isocitrate dehydrogenase [Bacillus subtilis]		86 78	1320
339	1 2	1489	1109	gi 1118003	[dihydroneopterin aldolase [Staphylococcus haemolyticus]	8	77 1 98	381
358	7 5	2124	3440	gi 1146219 	[28.2% of identity to the Escherichia coli GTP-binding protein Era; putative [Bacillus subtilis]		86 73	1317

Table (

| % ident | length (nt) # sim pha chain glycerol kinase (glpK) (EC 2.7.1.30) [Bacillus subtilis] pir|B45868|B45868 | glycerol kinase (EC 2.7.1.30) - Bacillus subtilis sp|P18157|GLPK_BACSU | GLYCEROL KINASE (EC 2.7.1.30) (ATP:GLYCEROL -PHOSPHOTRANSFERASE) Iglycerladehyde-3-phosphate dehydrogenase [Clostridium pasteurianum] pir18342541834254 glyceraldehyde-3-phosphate dehydrogenase (EC .2.1.12) ir|S10798|DEBSPF pyruvate dehydrogenase (lipoamide) (EC 1.2.4.1) |pyruvate dehydrogenase (lipoamide) [Bacillus stearothermophilus] 13-ketoacyl-acyl carrier protein reductase [Bacillus subtilis] [pyrimidine nucleoside transport protein [Bacillus subtilis] |phoP protein (put.); putative [Bacillus subtilis] |transmembrane protein [Saccharomyces cerevisiae] lasparaginyl-tRNA synthetase [Bacillus subtilis] ORTHININE AMINOTRANSFERASE [Bacillus subtilis] [glcB gene product [Staphylococcus carnosus] lipa-90d gene product [Bacillus subtilis] (thrSv) (EC 6.1.1.3) [Bacillus subtilis] [PET112-like protein [Bacillus subtilis] [unknown [Staphylococcus haemolyticus] |unknown (Staphylococcus haemolyticus) phoP gene product [Bacillus subtilis] |EpiF [Staphylococcus epidermidis] - Bacillus stearothermophilus [ATPase [Lactococcus lactis] [unknown [Bacillus subtilis] [unknown [Bacillus subtilis] Clostridium pasteurianum [YqkL [Bacillus subtilis] |YqfA [Bacillus subtilis] |YqhW [Bacillus subtilis] (GK) (GLYCEROKINASE) match gene name acession 1gi | 1303817 |gi|1176399 |gi|1146247 gi | 1303912 gi | 1303993 gil1022726 |gi|1022725 |gi|1502421 |gi|1408507 | 3475 | 5673 | 191|1072419 match |gi|1002911 |gi|1354211 | 6466 | 5252 |gi|1064807 1gi1467376 |gi|143328 gi|414014 gi | 296464 |gi|142992 |gi|143766 |gi|311924 1gi1467385 |gi|40056 |gi|40041 (nt) 1 240 1 162 Start 1 4287 (nt) 1 332 1 476 11 1 2 Contig |ORF | 1 2 ΩI ~ _ - 5 _

Table ?

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	10RF	Start (nt)	Stop (nt)	match acession	match gene name	mis as	* ident	length (nt)
1 205	115	8624	8106	gi 1044981		85	75	519
1 205	120	110928	110596	pir A02819 R5BS ribosomal		85	72	333
1 220	9	6490	6101	gi 48980	secA gene product (Bacillus subtilis)	85	99	390
231	4 -	1 4877	3159	gi 1002520	Muts [Bacillus subtilis]	85	70	1719
1 243	6	8013	8783	gi 414011		85	72	771
1 249	7 -	5894	3186	gi 1405454		85	73	2709
302		140	475	gi 40173 		80 S0	72	336
1 333	-	5445	1 2968	gi 442360	ClpC adenosine triphosphatase [Bacillus subtilis]	85	69	2478
364	9 -	1 6082	8196	gi 871784	Clp-like ATP-dependent protease binding subunit [Bos taurus]	85	89	2115
1 448	- 5	1992	1339	gi 405134	acetate kinase [Bacillus subtilis]	85	89	654
747		1251	853	(gi 1373157	orf-X; hypothetical protein; Method: conceptual translation supplied by author [Bacillus subtilis]	85	73	399
988	7 -	159	1 467	gi 541768		85	55	309
1 1089		1208	909	pir B47154 B471	pir B47154 B471 Signal recognition particle 54K chain homolog Ffh - Bacillus subtilis	85	71	603
1163		816	409	gi 304155 	diaminopimelate decarboxylase (Bacillus methanolicus) sp P41023 DCDA_BACMT DIAMINOPIMELATE DECARBOXYLASE (EC 4.1.1.20) DAP DECARBOXYLASE).	85	62	408
1 1924		1 487	1 251	911215098	excisionase (Bacteriophage 154a)	85	73	237
1 2932		977	1 390	gi 1041099		85	71	387
3030		e 	1 275	gi 42370 	pyruvate formate-lyase (AA 1-760) [Escherichia coli] ir S01788 S01788 formate C-acetyltransferase (EC 2.3.1.54) - cherichia coli	85	74	273
3111	·	1 595	1 299	gi 63568	limb deformity protein [Gallus gallus]	85	85	297
3778	-	1 630	316	gi 391840	beta-subunit of HDT [Pseudomonas fragi]	85	67	315
1 3835	-		1 387	gi 1204472	type I restriction enzyme ECOR124/3 I M protein [Haemophilus influenzae]	85	56	387
4042		۳ 	386	gi 18178 	formate acetyltransferase [Chlamydomonas reinhardtii] ir 524997 524997 formate C-acetyltransferase (EC 2.3.1.54) - lamydomonas reinhardtii	85	70	384
4053		1 35	340	gi 1204472	type I restriction enzyme ECOR124/3 I M protein [Haemophilus influenzae]	85	56	306
4108		7	181	gi 1072418	glcA gene product [Staphylococcus carnosus]	85	61	180
4300		1 575	330	gi 151932	[fructose enzyme II [Rhodobacter capsulatus]	85	59	246
-	1		1					

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	g IORF	Start (nt)	Stop (nt)	match acession	match gene name	e sim	% ident	length (nt)
1 4392		1 627	1 355	gi 1022725	unknown (Staphylococcus haemolyticus)	85	14	273
4408		- 2	1 235	gi 871784	Clp-like ATP-dependent protease binding subunit [Bos taurus]	85	62	234
1 4430		1 578	1 291	gi 1009366	Respiratory nitrate reductase [Bacillus subtilis]	85	89	288
4555		2	1 253	gi 450688	HadM gene of Ecopril gene product (Escherichia coli) pir(538477(538437 hadM protein - Escherichia coli pir(509629 509629 hypothetical protein A - Escherichia coli (SUB 40-520)	8 1 S 2 S	252	252
4611	·	481	1 242	gi 1256635	dihydroxy-acid dehydratase [Bacillus subtilis]	85	65	240
4	110	110061	110591	gi 46982	fosB gene product [Staphylococcus epidermidis]	84	1 89 1	531
- 13	5	1 1348	1172	gi 142450		84	56	177
1 16	4 -	1803	4652	gi 1277198	DNA repair protein [Deinococcus radiodurans]	84	67	2850
1 22	e -	1535	1128	gi 511069	Uref [Staphylococcus xylosus]	84	73	408
1 23	7	5055	5306	gi 603320	Yer082p [Saccharomyces cerevisiae]	84	61	252
53	11	111597	111145	gi 1303948	YqiW [Bacillus subtilis]	84	89	453
53	17	114059	112770	gi 142613 	branched chain alpha-keto acid dehydrogenase E2 [Bacillus subtilis] gi 1303944 BfmBB [Bacillus subtilis]	84	71	1290
10		1332	1 982	gi 46647	ORF (repE) [Staphylococcus aureus]	84	68	351
1 73	4	1 2512	4311	gi 142993	glycerol-3-phosphate dehydrogenase (glpb) (EC 1.1.99.5) [Bacillus ubtilis]	84	74	1800
86	7	4324	9609	gi 467427	methionyl-tRNA synthetase [Bacillus subtilis]	84	99	1773
100	6	9501	1 8680	gi 1340128	ORF1 [Staphylococcus aureus]	84	78	822
111	. –	1934	1 3208	gi 1237019	Srb (Bacillus subtilis]	84	1 69 1	1275
1 148	9	1 4720	1 5670	gi 467462	cysteine synthetase A [Bacillus subtilis]	84	1 69 1	951
152	4	2064	1 2456	gi 143377 	pyruvate decarboxylase (E-1) alpha subunit [Bacillus subtilis] pir B36718 DEBSPA pyruvate dehydrogenase (lipoamide) (EC 1.2.4.1) lpha chain - Bacillus subtilis	84	70	393
1 169	7	1 3634	3861	gi 1001342	hypothetical protein [Synechocystis sp.]	84	99	228
171	4	1 2992	1 2657	gi 517475	D-amino acid transaminase [Staphylococcus haemolyticus]	84	71	336
1 186	9	6941	6216	gi 467475	unknown (Bacillus subtilis)	84	102	726
1 205	6	6261	1 5692	gi 216340	[ORF for adenylate kinase [Bacillus subtilis]	84	71	570
1 224	7	915	1 1391	gi 288269	beta-fructofuranosidase [Staphylococcus xylosus]	84	1 0/	477

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

	1	1 +		111111111111111111111111111111111111111		+	+	+
Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
251	1	92	388	gi 1303790	Yqel [Bacillus subtilis]	84	65	297
282	e 	1526	2836	gi 143040 	glutamate-1-semialdehyde 2,1-aminotransferase (Bacillus subtilis) pir10427281042728 glutamate-1-semialdehyde 2,1-aminomutase (EC .4.3.8) - Bacillus subtilis	84	75	1311
307	2	3138	2959	gi 1070014	protein-dependent (Bacillus subtilis)	84	62	180
1 320	4	2343	1 4229	gi 143390	carbamyl phosphate synthetase [Bacillus subtilis]	84	70	1887
372		8	1 296	gi 1022725	unknown [Staphylococcus haemolyticus]	84	70	294
413	- 5	2201	1341	gi 1256146	YbbQ [Bacillus subtilis]	84	65	861
439	-	3	392	gi 1046173	osmotically inducible protein [Mycoplasma genitalium]	84	53	390
461	e 	1362	1 2270	gi 40211	Threonine synthase (thrC) (AA 1-352) [Bacillus subtilis] ir A25364 A25364 threonine synthase (EC 4.2.99.2) - Bacillus btilis	84	69	1 606
1 487	1	3	1 299	gi 1144531	integrin-like protein alpha Intlp [Candida albicans]	84	46	297
491	2	624	1 905	pir S08564 R3BS	Iribosomal protein S9 - Bacillus stearothermophilus	84	69	282
1 491	8	1 836	1033	pir S08564 R3BS	Iribosomal protein S9 - Bacillus stearothermophilus	84	77	198
548	-	3	341	1911431231	uracil permease (Bacillus caldolyticus)	84	74	339
128	7	2701	1748	gi 912445	DNA polymerase [Bacillus caldotenax]	84	89	954
1 769	1	3	1 257	[gi[1510953	cobalamin biosynthesis protein N [Methanococcus jannaschii]	84	38	255
954		308	156	gi 1405454	aconitase (Bacillus subtilis)	84	57	153
1 957		ε	395	gi 143402 	recombination protein (ttg start codon) [Bacillus subtilis] gi 1303923 RecN [Bacillus subtilis]	84	89	393
975	1	e .	452	gi 885934	[Clp8 [Synechococcus sp.]	84	70	450
1585		3	1 257	gi 510140	ligoendopeptidase F [Lactococcus lactis]	84	56	255
1 2954		3	323	gi 603769	HutU protein, urocanase [Bacillus subtilis]	84	73	321
1 2996		650	348	gi 18178 	formate acetyltransferase [Chlamydomonas reinhardtii] ir 524997 524997 formate C-acetyltransferase (EC 2.3.1.54) - lamydomonas reinhardtii	84	65	303
3766		737	375	gi 517205	67 kDa Myosin-crossreactive streptococcal antigen (Streptococcus yogenes)	84	72	363
4022		2	169	gi 1146206	glutamate dehydrogenase [Bacillus subtilis]	84	54	168
4058		1 620	312	gi 151932	[fructose enzyme II [Rhodobacter capsulatus]	84	71	309
4108	- 5	1 106	1 351	gi 1072418	glcA gene product [Staphylococcus carnosus]	84	177	246
	1 1 1 1 1			-				

Table

| % sim | % ident | length (nt) | pir|A35260|A35260 alcohol dehydrogenase (EC 1.1.1.1) I - Zymomonas obilis imidazoleglycerol-phosphate synthase (cyclase) [Methanococcus jannaschii] |glutamate synthase large subunit (EC 2.6.1.53) [Escherichia coli]
| pir/A29617|A29617 glutamate synthase (NADPH) (EC 1.4.1.13) large hain
| Escherichia coli |branched chain alpha-keto acid dehydrogenase E1-beta (Bacillus ubtilis) S. aureus - Putative coding regions of novel proteins similar to known proteins lalcohol dehydrogenase I (adhA) (EC 1.1.1.1) [Zymomonas mobilis] [lumazine synthase (b-subunit) [Bacillus amyloliquefaciens] |replication initiator protein [Listeria monocytogenes] linitiation protein of replicaton (Bacillus subtilis) ORF for L30 ribosomal protein (Bacillus subtilis) |fumarase (citG) (aa 1-462) (Bacillus subtilis] [phospho-beta-glucosidase [Bacillus subtilis] cysteinyl-tRNA synthetase (Bacillus subtilis) |HutU protein, urocanase (Bacillus subtilis) |urease accessory protein [Bacillus sp.] |GTP-binding protein [Bacillus subtilis] |ORF6 gene product [Bacillus subtilis] (ORF2 gene product [Bacillus subtilis] |unknown [Bacillus subtilis] |ORF75 (Bacillus subtilis) |ORFU [Lactococcus lactis] |YqhK [Bacillus subtilis] |YqhM [Bacillus subtilis] orf4 [Bacillus subtilis] |kdpB [Escherichia coli] | match gene name | aldolyticus acession | 3 | 3119 | 2274 | gi | 1303894 |gi|1212730 1gi|1212776 |gi|1510488 |gi|1039479 match |gi|146208 1gi | 603769 |14722 |13745 |gi|142612 |gi|155571 |gi|506381 | 5749 | 6696 | 19i1903307 lgi|520844 |gi|467419 1gi | 508979 | 4 | 2043 | 1576 |gi|393297 |gi|143132 1gi | 634107 |14 |14148 |14546 |gi|580902 1gi1289284 |gi|216337 |gi|467391 |11070 |gi|49316 | 1864 | 1523 |gi|44095 |gi|39844 1 2467 1 2742 | 4 | 1421 | 2743 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 1 7607 1 694 1 638 1 352 1 9880 1 4 | 3933 1 4 1 3209 1 1 985 ж — 1 1 2 _ ----~ <u>س</u> 1 2

Table?

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig l	10RF	Start (nt)	Stop (nt)	match acession	match gene name	+	* ident	length (nt)
387	† - ·	3	662	 gi 806281	DNA polymerase I [Bacillus stearothermophilus]	++	1 02	099
527	2	916	1566	gi 396259	protease [Staphylococcus epidermidis]	83	1 19	651
533		355	179	gi 142455 	nalanine dehydrogenase [EC 1.4.1.1) [Bacillus stearothermophilus] pir B34261 B34261 alanine dehydrogenase (EC 1.4.1.1) - Bacillus tearothermophilus	8 83	99	177
536	4	1617	1438	gi 143366 	adenylosuccinate lyase (PUR-B) (Bacillus subtilis] pir C29326 WZBSDS adenylosuccinate lyase (EC 4.3.2.2) - Bacillus ubtilis	83	67	180
652		2	829	gi 520753	DNA topoisomerase I [Bacillus subtilis]	83 1	72	828
774	2 -	200	361	gi 1522665	[M. jannaschii predicted coding region MJECL28 [Methanococcus jannaschii]	83	58	162
1 68		120	296	gi 1064807	ORTHININE AMINOTRANSFERASE [Bacillus subtilis]	83	1 94	177
1213	7	6	491	gi 289288	lexA [Bacillus subtilis]	83	1 19	489
2529		296	150	gi 143786 	tryptophanyl-tRNA synthetase (EC 6.1.1.2) [Bacillus subtilis] pir JT0481 YWBS tryptophantRNA ligase (EC 6.1.1.2) - Bacillus ubtilis	83	69	147
2973		649	326	gi 1109687	ProZ [Bacillus subtilis]	83	58	324
3009	-	728	366	gi 882532	ORF_0294 [Escherichia coli]	83 -	65 1	363
3035	7	45	305	gi 950062 	hypothetical yeast protein 1 [Mycoplasma capricolum] pir S48578 S48578 hypothetical protein - Mycoplasma capricolum SGC3) (fragment)	83	59	261
3906		67	309	gi 1353197	[thioredoxin reductase [Eubacterium acidaminophilum]	83	61 +	243
4458	-	540	271	gi 397526	clumping factor (Staphylococcus aureus)	83	1 87	270
4570	1	444	223	gi 1022726	unknown [Staphylococcus haemolyticus]	83	74	222
4654	1	97	261	gi 1072419	glcB gene product [Staphylococcus carnosus]	83	1 62	165
16	7	295	1191	1911153854	luvs402 protein (Streptococcus pneumoniae)	82	1 29	897
16	e e	1193	1798	gi 153854	uvs402 protein [Streptococcus pneumoniae]	82	1 07	909
38	112	9644	8724	gi 1204400	N-acetylneuraminate lyase [Haemophilus influenzae]	82	58	921
42	4.	886	2019	gi 841192	catalase [Bacteroides fragilis]	82	1 07	1032
51	9	2590	3489	gi 143607	sporulation protein (Bacillus subtilis)	82	1 69	006
56	11	12270	13925	gi 39431	oligo-1,6-glucosidase [Bacillus cereus]	82	1 09	1656
56	115 1	17673	18014	gi 467410	unknown [Bacillus subtilis]	82 +	1 99	342
61	2	881	3313	gi 143148	transfer RNA-Leu synthetase [Bacillus subtilis]	82	1 07	2433
+						+ +		! !

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig ORF	ORF	Start (nt)	Stop (nt)	match acession	match gene name	mis s	% ident	length (nt)
8		9162	11318	gi 48240 	elongation factor G (AA 1-691) [Thermus aquaticus thermophilus] ir S15928 EFTWG translation elongation factor G - Thermus aquaticus p P1355 EFG_THETH ELONGATION FACTOR G (EF-G).	82	64	2157
85	7	5470	3260	gi 143369	phosphoribosylformyl glycinamidine synthetase II (PUR-Q) [Bacillus ubtilis]	82	99	2211
102	9	3662	1 5380	gi 1256635	dihydroxy-acid dehydratase [Bacillus subtilis]	82	1 59	1719
117	4	3242	1 3493	pir A47154 A471	orf1 5' of Ffh - Bacillus subtilis	82	53	252
128	9	4377	5933	gi 460258	phosphoglycerate mutase [Bacillus subtilis]	82	1 99	1557
129	8	1229	2182	gi 403373 	glycerophosphoryl diester phosphodiesterase [Bacillus subtilis] pir[537251 537251 glycerophosphoryl diester phosphodiesterase - acillus subtilis	82	62	954
170	-	7	1441	gi 1377831	unknown [Bacillus subtilis]	82	67	1440
177	-	۳ ا	1094	gi 467386	(thiophen and furan oxidation [Bacillus subtilis]	82	65	1092
184	4	3572	4039	gi 153566	ORF (19K protein) Enterococcus faecalis]	82	59	468
189	8	4455	1 4225	gi 1001878	(CspL protein [Listeria monocytogenes]	82	73	231
1 206	119	121366	120707	gi 473916	Ilipopeptide antibiotics iturin A (Bacillus subtilis) sp P39144 LP14 BACSU LIPOPEPTIDE ANTIBIOTICS ITURIN A AND SURFACTIN IOSYNTHESIS PROTEIN.	82	50	099
1 221	7	805	1722	gi 517205	167 kDa Myosin-crossreactive streptococcal antigen [Streptococcus yogenes]	82	63	918
223	4	3866	3651	gi 439619 	[Salmonella typhimurium IS200 insertion sequence from SARA17, artial.], gene product [Salmonella typhimurium]	82	69	216
1 260	8	5207	1 4296	gi 1161381		82	61	912
315	3	4864	2855	gi 143397	quinol oxidase [Bacillus subtilis]	82	67	2010
321	110	8520	7945	gi 142981 	ORF5; This ORF includes a region (aa23-103) containing a potential ron- sulphur centre homologous to a region of Rhodospirillum rubrum nd Chromatium vinosum; putative [Bacillus stearothermophilus] pir[PQ0299[PQ0299 hypothetical protein 5 (gldA 3' region) -	82	62	576
331	e -	1055	1342	gi 436574	ribosomal protein L1 (Bacillus subtilis	82	71	288
370	2	262	618	gi 1303793	YqeL [Bacillus subtilis]	82	59	357
1 404	4	3053	1 4024	gi 1303821	YqfE [Bacillus subtilis]	82	89	972
1 405	4	4440	1 3073	gi 1303913	ΥqhX [Bacillus subtilis]	82	1 69	1368
436	m 	4096	2864	gi 149521 	tryptophan synthase beta subunit [Lactococcus lactis] pir(835129 835129 tryptophan synthase (EC 4.2.1.20) beta chain - actococcus lactis subsp. lactis	88 2	67	1233

Table ?

S. aureus - Putative coding regions of novel proteins similar to known proteins

				-		1	4	1
Contig ORF	9 JORF	Start (nt)	Stop (nt)	match acession	match gene name	a sim	* ident	length (nt)
4	4	3394	1 2573	gi 142952	glyceraldehyde-3-phosphate dehydrogenase [Bacillus tearothermophilus]	82	1 29	822
1 444	112	110415	111227	gi 1204354	spore germination and vegetative growth protein [Haemophilus influenzae]	82	67	813
1 446		e - ·	191	gi 143387	aspartate transcarbamylase [Bacillus subtilis]	82	1 99	189
462	m 	1007	1210	gi 142521 	decoxyribodipyrimidine photolyase [Bacillus subtilis] pir[A37192[A37192 uvrB protein - Bacillus subtilis sp P14951 UVRC_BACSU EXCINUCLEASE ABC SUBUNIT C.	85	64	204
1 537		1 1560	1 784	gi 853767	UDP-N-acetylglucosamine 1-carboxyvinyltransferase [Bacillus ubtilis]	82	61	177
1 680	2 -	407	1 700	gi 426472	secE gene product (Staphylococcus carnosus]	82	69	294
724	7	1 565	386	gi 143373	$Iphosphoribosyl\ aminoimidazole\ carboxy\ formyl\ ormyltransferase/inosine\\ Imponophosphate\ cyclohydrolase\ (PUR-H(J))\ Bacillus\ subtilis]$	82	89	180
1 763	1	1 422	1 213	gi 467458	cell division protein (Bacillus subtilis)	82	35	210
818		1 564	1 283	gi 1064787	[function unknown [Bacillus subtilis]	82	1 69	282
828		175	1176	gi 143043	uroporphyrinogen decarboxylase [Bacillus subtilis] pir B47045 B47045 uroporphyrinogen decarboxylase (EC 4.1.1.37) - acillus subtilis	82	71	1002
895		m 	599	gi 1027507	ATP binding protein (Borrelia burgdorferi)	82	72	597
686	1	10	399	gi 143795	transfer RNA-Tyr synthetase [Bacillus subtilis]	82	09	390
961			306	gi 577647	gamma-hemolysin [Staphylococcus aureus]	82	69	306
1 1192		1 307	155	gi 146974	HH3-dependent NAD synthetase [Escherichia coli	82	71	153
1317	-	49	375	gi 407908		82	72	327
1341			150	gi 39962 	ribosomal protein L35 (AA 1-66) [Bacillus stearothermophilus} ir S05347 R5BS35 ribosomal protein L35 - Bacillus earothermophilus	82	89	150
1 2990	~	1 567	349	gi 534855	ATPase subunit epsilon [Bacillus stearothermophilus] splP42009 ATPE_BACST ATP SYNTHASE EPSILON CHAIN (EC 3.6.1.34).	85	47	219
3024	- -	45	1 224	gi 467402	unknown (Bacillus subtilis)	82	64	180
3045	-	1 276	139	gi 467335	[ribosomal protein L9 (Bacillus subtilis]	82	09	138
3045	- 5	558	400	gi 467335	ribosomal protein L9 [Bacillus subtilis]	82	82	159
3091		474	1 238	gi 499335	secA protein [Staphylococcus carnosus]	82	78	237
3107	-	416	210	gi 546918 	orfY 3' of comK [Bacillus subtilis, E26, Peptide Partial, 140 aa] pir S43612 S43612 hypothetical protein Y - Bacillus subtilis sp P40398 YHXD_BACSU HYPOTHETICAL PROTEIN IN COMK 3'REGION (ORFY) FRAGMENT).	82	64	207
	+			,,,,,,,,,,,,,,,,,,,,,				

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

+	+	+	+					
Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
4332		7	319	gi 42086 	Initrate reductase alpha subunit (Escherichia coli) p/P09152 NARG_ECOLI RESPIRATORY NITRATE REDUCTASE 1 ALPHA CHAIN (EC 7.99.4). (SUB 2-1247)	82	75	318
23	<u>م</u>	3275	2574	gi 1199573	spsB [Sphingomonas sp.]	81	1 64	702
42		638	321	gi 466778	lysine specific permease [Escherichia coli]	81	59	318
1 48		1 4051	1 4350	gi 1045937	IM. genitalium predicted coding region MG246 [Mycoplasma genitalium]	81	62	300
51	4	1578	1 2579	pir S16649 S166	ldciAC protein - Bacillus subtilis	81	55	1002
53	- 5	364	1494	gi 1303961	ΥqjJ [Bacillus subtilis]	81	67	1131
53	80	9419	7971	gi 146930	6-phosphogluconate dehydrogenase [Escherichia coli]	81	99	1449
54	ъ ъ	110757	10119	gi 143016	permease {Bacillus subtilis}	81	65	629
54	110	113360	111786	gi 143015	gluconate kinase [Bacillus subtilis]	81	64	1575
57	117	113983	13366	pir A25805 A258	L-lactate dehydrogenase (EC 1.1.1.27) - Bacillus subtilis	81	74	618
81	- 5	1 2708	1 2217	lgi 1222302	NifU-related protein [Haemophilus influenzae]	81	54	492
98		1 745	374	gi 414017	lipa-93d gene product [Bacillus subtilis]	81	1 04	372
103	9	6438	4861	gi 971342 	Initrate reductase beta subunit (Bacillus subtilis) sp P42176 NARH_BACSU NITRATE REDUCTASE BETA CHAIN (EC 1.7.99.4).	81	64	1578
120	115	110845	112338	gi 1524392	GbsA [Bacillus subtilis]	81	1 69	1494
128	2	3676	4413	gi 143319	triose phosphate isomerase [Bacillus megaterium]	81	64	738
131	6	110308	9280	gi 299163	alanine dehydrogenase (Bacillus subtilis)	81	89	1029
143	9	8809	5471	gi 439619 	[Salmonella typhimurium IS200 insertion sequence from SARA17, artial.], gene product [Salmonella typhimurium]	81	61	618
169		43	825	gi 897795	130s ribosomal protein [Pediococcus acidilactici] sp1P49668 RS2_PEDAC 30S RIBOSOMAL PROTEIN S2.	81	65	783
230		450	1 226	gi 1125826 	short region of weak similarity to tyrosine-protein kinase receptors in a fibronectin type III-like domain [Caenorhabditis elegans]	81	54	225
233	2	1 2000	1 2677	gi 467404	unknown [Bacillus subtilis]	81	63	678
241	~	3081	2149	gi 16510 	succinateCoA ligase (GDP-forming) [Arabidopsis thaliana] ir 330579 530579 succinateCoA ligase (GDP-forming) (EC 6.2.1.4) pha chain - Arabidopsis thaliana (fragment)	81	69	933
256		. – .	981	pir S09411 S094	spoIIE protein - Bacillus subtilis	81	65	981
1 259	. —	3 3752	1 2691	sp P28367 RF2_B	PROBABLE PEPTIDE CHAIN RELEASE FACTOR 2 (RF-2) (FRAGMENT).	81	65	1062

Table

1308 300 240 309 1068 573 561 996 174 903 609 237 561 345 399 150 189 222 237 324 399 | % sim | % ident | length 19 69 63 64 09 67 63 65 65 65 57 26 71 89 72 43 43 62 28 81 81 81 81 [methylated-DNA--protein-cysteine methyltransferase [Haemophilus influenzae] |synergohymenotropic toxin [Staphylococcus intermedius] pir|S44944|S44944 [{Salmonella typhimurium IS200 insertion sequence from SARA17, artial.], |L-glutamine-D-fructose-6-phosphate amidotransferase (Bacillus ubtilis] IH. influenzae predicted coding region HI0594 [Haemophilus influenzae] |Manganese superoxide dismutase [Bacillus caldotenax] ir|S22053|S22053 | superoxide dismutase (EC 1.15.1.1) (Mn) - Bacillus Idotenax IM. genitalium predicted coding region MG423 [Mycoplasma genitalium] |pir|C37083|C370 |hypothetical protein II (ompH 3' region) - Salmonella typhimurium |dipeptide transport ATP-binding protein [Haemophilus influenzae] |helicase [Autographa californica nuclear polyhedrosis virus] |mevalonate pyrophosphate decarboxylase [Rattus norvegicus] synergohymenotropic toxin - Staphylococcus ntermedius [aspartyl-tRNA synthetase [Haemophilus influenzae] [adenylosuccinate synthetase [Bacillus subtilis] [protein kinase PknB [Mycobacterium leprae] gene product [Salmonella typhimurium] |pir|S09411|S094 |spoIIIE protein - Bacillus subtilis (transketolase [Bacillus subtilis] sp/P24307/V143_NPVAC HELICASE. |resolvase [Transposon Tn917] [unknown [Bacillus subtilis] |ORF f388 [Escherichia coli] (ATPase [Lactococcus lactis] |YqhU [Bacillus subtilis] [gueA [Escherichia coli] |yeeD [Escherichia coli] |Bex [Bacillus subtilis] match gene name |gi|1204844 |gi|1204570 1gi | 1204652 1gi | 1262360 lgi|1303902 1gi11405446 |gi|1205429 |gi|1046138 |gi|1322245 |gi|467416 |gi|537137 |gi|467328 igi|154961 | 6154 | | gil606745 gi1487686 |gi|559164 |gi|439619 |gi|296464 |gi|147485 |gi|405955 |gi|39453 5889 1064 Contig |ORF | Start | Stop ID |ID | (nt) | (nt) 1406 1 5889 1342 1 2471 735 603 1 430 1 400 241 189 239 401 401 449 325 1 1728 6251 5653 1466 1 5590 11137 1034 1 5252 1404 1 1707 1727 1105 66 98 43 504 798 670 9 1 475 252 1 3940 | 1 | 3 _ --5 _ -<u>-</u> 7 7 7 5 7 _ 6 ~ 7 ~ 1035 3371 1280 3908 480 296 302 317 343 360 372 392 404 754 953 364

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	-+	match gene name	8 sim	* ident	length (nt)
1 3954		1	1 318	gi 1224069	amidase [Moraxella catarrhalis]	81	1 89	318
4049		337	170	gi 603768	Huti protein, imidazolone-5-propionate hydrolase [Bacillus subtilis] gi 603768 Huti protein, imidazolone-5-propionate hydrolase Bacillus subtilis}	81	89	168
4209			324	gi 403373	(glycerophosphoryl diester phosphodiesterase [Bacillus subtilis] pir(S37251 S37251 glycerophosphoryl diester phosphodiesterase - acillus subtilis	81	28	324
4371		627	322	gi 216677	<pre>lindolepyruvate decarboxylase [Enterobacter cloacae] pir(S16013 S16013 lindolepyruvate decarboxylase (EC 4.1.1) - nterobacter cloacae</pre>	81	72	306
4387		1 19	1 228	gi 460689	[TVG [Thermoactinomyces vulgaris]	81	59	210
4391	-	1 581	306	gi 1524193	unknown [Mycobacterium tuberculosis]	81	67 1	276
1 4425		e -	341	gi 143015	[gluconate kinase [Bacillus subtilis]	81	1 99	339
6	1	1593	1 847	gi 1064786	function unknown [Bacillus subtilis]	1 08	62	747
17		544	311	gi 559164 	helicase [Autographa californica nuclear polyhedrosis virus] sp P24307 V143 NPVAC HELICASE.	08	40	234
45	7	1159	1 2448	gi 1109684	ProV {Bacillus subtilis}	08	63	1290
1 45	- 2	1 4032	1 4733	gi 1109687	ProZ [Bacillus subtilis]	08	55	702
54	ъ - — -	110266	9502	gi 563952	gluconate permease [Bacillus licheniformis]	1 08 1	62	765
1 62	112	1 8852	7545	gi 854655	Na/H antiporter system [Bacillus alcalophilus]	1 88 1	62	1308
1 62	114	1 8087	8683	gi 559713	ORF [Homo sapiens]	08	1 89	597
1 67	116	113781	114122	gi 305002	ORF_f356 [Escherichia coli]	08	65	342
07	113	111495	110296	gi 1303995	YqkN [Bacillus subtilis]	80	64	1200
86	6	6336	7130	gi 467428	unknown [Bacillus subtilis]	1 80 1	1 89	795
86	110	1 7294	1 7833	gi 467430	unknown [Bacillus subtilis]	1 80 1	64	540
86	=	1 7820	1 8737	gi 467431	high level kasgamycin resistance [Bacillus subtilis]	08	61	918
109	116	114154	114813	gi 580875	ipa-57d gene product [Bacillus subtilis]	80	63	1 099
112	115	114294	116636	gi 1072361	pyruvate-formate-lyase [Clostridium pasteurianum]	08	65	2343
139	. –	1448	1 726	gi 506699	CapC [Staphylococcus aureus]	1 08	58	723
139	- 1	1 2179	1 1448	gi 506698	CapB [Staphylococcus aureus]	1 80 1	59	732
174		3271	1 2870	gi 1146242	aspartate 1-decarboxylase [Bacillus subtilis]	08	61	402

Table

| % ident | length | % sim Matches PS00017: ATP GTP A and PS00301: EFACTOR GTP; similar to longation |valy1-tRNA synthetase [Bacillus stearothermophilus] sp|P11931|SYV_BACST | factor G, TetM/TetO tetracycline-resistance proteins Escherichia coli) |proton glutamate symport protein [Bacillus stearothermophilus] | pir{S26247|S26247 glutamate/aspartate transport protein - Bacillus | tearothermophilus S. aureus - Putative coding regions of novel proteins similar to known proteins VALYL-TRNA SYNTHETASE (EC 6.1.1.9) VALINE--TRNA LIGASE) (VALRS) |formyl-tetrahydrofolate synthetase (Streptococcus mutans) |ribosomal protein L4 [Bacillus stearothermophilus] |beta-fructofuranosidase [Staphylococcus xylosus] |putative ATP binding subunit (Bacillus subtilis) |glutamyl-tRNA synthetase [Bacillus subtilis] |85-kDa surface antigen [Trypanosoma cruzi] lacuC gene product [Staphylococcus xylosus] lacetyl-CoA synthetase [Bacillus subtilis] [hypothetical protein [Bacillus subtilis] |outer capsid protein [Rotavirus sp.] unknown [Mycobacterium tuberculosis] |function umknown [Bacillus subtilis] |function umknown [Bacillus subtilis] spec [Streptococcus pyogenes] |putative (Bacillus subtilis) [enolase [Bacillus subtilis] |unknown [Bacillus subtilis] IORF_o470 [Escherichia coli] |unknown [Bacillus subtilis] |unknown [Bacillus subtilis] ORFX7 [Bacillus subtilis] |YneF [Bacillus subtilis] acession |gi|1405448 |gi|1146183 |gi|1483545 |gi|1103865 |gi|1177686 |gi|1064791 |gi|1064791 |gi|161953 |gi|143000 |gi|467418 |gi|310325 1gi1537049 1gi1467448 |gi|304976 |gi|529754 1gi | 289282 |gi|143797 | 5 | 2724 | 2461 |gi|709993 |gi|348053 |gi|460259 |gi|410131 1gi140103 Stop (nt) 1 2255 1 2842 1 2240 1 7006 1 636 (nt) 1 2102 1 1 1268 Contig | ORF | <u>ش</u> _ ~ Ŋ

|HMG-CoA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir|A44756|A44756 167 kDa Myosin-crossreactive streptococcal antigen [Streptococcus yogenes] |large subunit of NADH-dependent glutamate synthase (Plectonema boryanum) large subunit of NADH-dependent glutamate synthase (Plectonema boryanum gi|603768 Hutl protein, imidazolone-5-propionate hydrolase Bacillus |Hut1 protein, imidazolone-5-propionate hydrolase (Bacillus subtilis) |HutI protein, imidazolone-5-propionate hydrolase [Bacillus subtilis] gil603768 HutI protein, imidazolone-5-propionate hydrolase Bacillus | hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp. |leukotoxin secretion ATP-binding protein [Haemophilus influenzae] S. aureus - Putative coding regions of novel proteins similar to known proteins |coenzyme PQQ synthesis protein III [Methanococcus jannaschii] fructose-permease IIBC component [Haemophilus influenzae] [photosystem I core protein B [Synechococcus vulcanus] |pir|A61072|EPSG |gallidermin precursor - Staphylococcus gallinarum Respiratory nitrate reductase [Bacillus subtilis] |UDP-glucose pyrophosphorylase [Bacillus subtilis] (2-nitropropane dioxygenase [Williopsis saturnus] [polynucleotide phosphorylase [Bacillus subtilis] [NADH-glutamate synthase [Medicago sativa] [deoxyribose aldolase [Mycoplasma hominis] | 6 | 5273 | 5515 |pir|A36728|A367 |acyl carrier protein - Rhizobium meliloti ORFX17 [Bacillus subtilis] orf6 [Lactobacillus sake] |IIGlc [Bacillus subtilis] [CodY [Bacillus subtilis] |dpj [Escherichia coli] |vp2 [Marburg virus] match gene name subtilis] | subtilis] acession 1gi | 1511600 |gi|1204696 |gi|1009366 |qi|1370207 |gi|1339950 gi | 1205301 |gi|1184680 |gi|1339950 |gi|517205 |gi|410141 gi | 147556 |gi|151259 |gi|217130 |gi|415855 |gi|166412 |gi|289287 |gi|603768 |gi|535351 |gi|603768 gi | 551531 1gi160632 |gi|39956 1 275 Contig |ORF | Start | ID |ID | (nt) | 1 5998 1 17 1 294 1 798 1 680 1 285 1 692 - 5 1 - 3 - 1 ---~ 4030 | 1

Table?

S. aureus - Putative coding regions of novel proteins similar to known proteins

	1	1111111111				+	+	+
Contig	ORF IID	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
1 59	7	1173	1 1424	gi 147923	threonine dehydratase 2 (EC 4.2.1.16) [Escherichia coli]	1 61	75 †	252
09		 	1 204	gi 666115	orfl upstream of glucose kinase (Staphylococcus xylosus) pir/S52351/S52351 hypothetical protein 1 - Staphylococcus xylosus	79	1 09	204
81		3002	1 1590	gi 466882	pps1; B1496_C2_189 [Mycobacterium leprae]	1 67	64	1413
1 85		1 7023	1 6505	gi 143364	phosphoribosyl aminoimidazole carboxylase I (PUR-E) [Bacillus ubtilis]	1 62	1 09	519
68	φ	1 5660	4554	gi 144906	product homologous to E.coli thioredoxin reductase: J.Biol.Chem. 1988) 263:9015-9019, and to F52a protein of alkyl hydroperoxide eductase from 1 S.typhimurium: J.Biol.Chem. (1990) 265:10535-10540; pen reading frame A [Clostridium pasteurianum]	9 6	35	1107
102	=	1 7489	8571	gi 143093 	ketol-acid reductoisomerase (Bacillus subtilis) sp[P37253]ILVC BACSU KETOL- ACID REDUCTOISOMERASE (EC 1.1.1.86) ACETOHYDROXY-ACID ISOMEROREDUCTASE) (ALPHA-KETO-BETA-HYDROXYLACIL EDUCTOISOMERASE).	79	64	1083
1 102	114	111190	112563	gi 149428	putative [Lactococcus lactis]	1 62	65	1374
1 127	6	17792	9372	gi 458688	PrfC/RF3 [Dichelobacter nodosus]	1 62	1 89	1581
139	<u>س</u>	1 2540	1983	gi 506697	CapA [Staphylococcus aureus]	1 62	55	558
144	7	1644	1156	gi 1498296	peptide methionine sulfoxide reductase (Streptococcus pneumoniae)	1 61	47	489
148	7	529	1098	gi 467457 	hypoxanthine-guanine phosphoribosyltransferase [Bacillus subtilis] gil467457 hypoxanthine-guanine phosphoribosyltransferase [Bacillus ubtilis]	79	59	570
150		1 965	1 591	gi 755602	unknown [Bacillus subtilis]	1 62	61	375
176		1039	587	gi 297874 	fructose-bisphosphate aldolase (Staphylococcus carnosus) pir1A499431A49943 fructose-bisphosphate aldolase (EC 4.1.2.13) - taphylococcus carnosus (strain TM300)	79	65	453
186	7	7584	6874	gi 1314298 	ORF5; putative Sms protein; similar to Sms proteins from Haemophilus influenzae and Escherichia coli [Listeria monocytogenes]	79	64	711
205	116	1 8887	8498	gi 1044980	ribosomal protein L18 [Bacillus subtilis]	1 62	70	390
211		1	1 519	gi 1303994	YqkM [Bacillus subtilis]	1 62	62	519
223	1 2	4183	1 2801	lgi1488430	alcohol dehydrogenase 2 [Entamoeba histolytica]	1 62	1 09	1383
243	8	8915	1 7896	gi 580883	ipa-88d gene product (Bacillus subtilis)	1 61	1 09	1020
279	4	3721	1 4329	gi 413930	lpa-6d gene product [Bacillus subtilis]	1 61	1 65	609
300	7	11	1393	lgi1403372	glycerol 3-phosphate permease [Bacillus subtilis]	1 62	62	1383
307	m — — -	1 2930	1935	gi 950062	hypothetical yeast protein 1 [Mycoplasma capricolum] pir S48578 S48578 hypothetical protein - Mycoplasma capricolum SGC3) (fragment)	79	09	966
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,								

Table

291 1221 576 504 351 624 420 579 300 399 156 384 285 210 183 312 219 474 267 243 297 282 333 | % ident | length (nt) 71 29 99 48 61 62 62 62 61 28 72 61 59 61 09 55 59 63 63 21 64 65 54 79 79 79 8 sim 79 79 79 79 79 79 79 19 79 |pir|C36889|C368 |leuB protein, inactive - Lactococcus lactis subsp. lactis (strain IL1403) [large subunit of NADH-dependent glutamate synthase [Plectonema boryanum] llarge subunit of NADH-dependent glutamate synthase (Plectonema boryanum larginine permease substrate-binding subunit [Listeria monocytogenes] 12-oxoglutarate dehydrogenase (odhA; EC 1.2.4.2) (Bacillus subtilis) |Clp-like ATP-dependent protease binding subunit [Bos taurus] pir|S52915|S529 |nitrate reductase alpha chain - Bacillus subtilis (fragment) [glycine betaine transporter OpuD [Bacillus subtilis] [Respiratory nitrate reductase [Bacillus subtilis] [3-phosphoglycerate kinase [Thermotoga maritima] pyruvate formate-lyase [Haemophilus influenzae] [nitrite reductase (nirB) [Bacillus subtilis] |orf-1; novel antigen [Staphylococcus aureus] lhypD gene product [Clostridium perfringens] |transport protein [Salmonella typhimurium] |Ribosomal Protein L10 [Bacillus subtilis] |deoxyribose aldolase [Mycoplasma hominis] [function unknown [Bacillus subtilis] [unknown [Mycobacterium tuberculosis] [gluconate kinase [Bacillus subtilis] |P47K [Pseudomonas chlororaphis] 6.3.3.1) - Bacillus subtilis [putative [Lactococcus lactis] [putative [Bacillus subtilis] [NisF [Lactococcus lactis] lorf6 (Lactobacillus sake) acession |gi|1064795 |gi|1407784 |gi|1204436 |gi|1149662 gi | 1339950 gi|1339950 |gi|1009366 |gi|1524193 |gi|1370207 |gi|1524397 |gi|871784 |gi|216854 |gi|143177 |gi|786163 |gi|805090 |gi|476160 |gi|149429 |gi|143267 gi|710020 gi | 450686 |gi|415855 |gi|143015 |gi|143371 110106 | 8886 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 1124 1 3987 352 374 1 421 314 476 1 312 341 401 202 385 286 411 270 332 326 401 14 | 3448 1153 1 2457 2663 4700 621 702 1 75 1 1 357 1 768 1 570 1 1 514 1 1 536 12 | 159 1 1 1 701 1 1 608 m 607 11 1 3 - 9 _ <u>س</u> œ — -<u>س</u> 2946 2937 2940 2999 3064 3126 3345 3836 2929 3022 3083 3181 816 352 481 683

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORE IID	Start Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
1 4501		1 406	1 209	gi 1022726	unknown [Staphylococcus haemolyticus]	1 62	99	198
1 4612	-	2	1 238	gi 460689		79	58	237
	-	7	1213	gi 520753	DNA topoisomerase I [Bacillus subtilis]	78	64	1212
	7	2266	1220	gi 216151	DNA polymerase (gene L; ttg start codon) [Bacteriophage SPO2] gil579197 SPO2 DNA polymerase (aa 1-648) [Bacteriophage SPO2] pir A21498 DJBPS2 DNA- directed DNA polymerase (EC 2.7.7.7) - phage PO2	187	12	1047
6	2	1340	1 1089	gi 1064787	function unknown [Bacillus subtilis]	78	57	252
32	8	1 6803	1 7702	gi 146974	NH3-dependent NAD synthetase [Escherichia coli]	78	63	006
1 36	4	1 2941	3138	gi 290503	glutamate permease [Escherichia coli]	78	53	198
1 53	115	117684	116221	gi 1303941	YqiV [Bacillus subtilis]	78	58	1464
1 57	114	110520	112067	gi 1072418	glcA gene product [Staphylococcus carnosus]	78	65	1548
99 1	7	62.98	1 5812	gi 1212729	YqhJ [Bacillus subtilis]	78	19	1 286
1 67	4	1 4029	1 4376	gi 466612	InikA [Escherichia coli]	78	7.1	348
91	6	110058	110942	gi 467380	stage 0 sporultion (Bacillus subtilis)	78	50	885
1 102	112	8574	110130	gi 149426	putative [Lactococcus lactis]	78	61	1557
112	9	3540	1 4463	gi 854234	lcymG gene product (Klebsiella oxytoca)	78	56	924
124	2	1888	1061	gi 405622	unknown [Bacillus subtilis]	78	1 09	828
1 130		1 1805	1 2260	gi 1256636	putative [Bacillus subtilis]	78	71	456
133	. –	1 751	1377	gi 168060	llamB [Emericella nidulans]	78	59	375
1 166	4	7125	6163	gi 451216	Mannosephosphate Isomerase [Streptococcus mutans]	78	63	963
1 186		1586	1 795	gi 289284	cysteinyl-tRNA synthetase [Bacillus subtilis]	78	63	792
1 195	4	2749	2315	gi 1353874	unknown [Rhodobacter capsulatus]	78	58	435
199	e 	4279	3623	gi 143525 	succinate dehydrogenase cytochrome b-558 subunit (Bacillus subtilis)	78	57	657
199	4	7209	5557	gi 142521	deoxyribodipyrimidine photolyase [Bacillus subtilis] pir[A371921A37192 uvrB protein - Bacillus subtilis sp P14951 UVRC_BACSU EXCINUCLEASE ABC SUBUNIT C.	78	62	1653
223	m 	3831	1 3523	gi 439596 	[Escherichia coli IS200 insertion sequence from ECOR63, partial.], ene	78	47	309
· -	-	1	-					

Table?

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig ORF ID ID 299 4	(F Start	t Stop	match	match gene name	# sim +	% ident	length
					1 1 1 1 1 1 1		· · · · · ·
	1865	1 2149	gi 467439	temperature sensitive cell division [Bacillus subtilis]	78	62	285
321 9	1 7734	7315	gi 142979	IORF3 is homologous to an ORF downstream of the spoT gene of E.coli, RF3 [Bacillus stearothermophilus]	78	55	420
352 4	1 3714	3944	gi 349050	lactin 1 [Pneumocystis carinii]	78	42	231
352 5	1 7592	- 6093	gi 903587	NADH dehydrogenase subunit 5 (Bacillus subtilis) sp P9975 NDHF BACSU NADH DEHYDROGENASE SUBUNIT 5 (EC 1.6.5.3) NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5).	78		1500
376 1	- 5	1 583	gi 551693	dethiobiotin synthase [Bacillus sphaericus]	78	34	582
424 2	1595	1768	gi 1524117	alpha-acetolactate decarboxylase [Lactococcus lactis]	78	99	174
450 1	1914	886	lgi 1030068	<pre>NAD(P)H oxidoreductase, isoflavone reductase homologue [Solanum tuberosum]</pre>	78	63	927
558 1	1 762	1 562	gi 1511588	bifunctional protein [Methanococcus jannaschii]	78	09	201
670 3	3 1152	1589		unknown [Bacillus subtilis]	78	64	438
714 1	1 64	1 732	gi 143460	137 kd minor sigma factor (rpoF, sigB; ttg start codon) [Bacillus ubtilis]	78	57	699
814 1	8	1 368	gi 1377833	unknown [Bacillus subtilis]	78	59	366
981 1	1381	1 692	gi 143802	GerC2 [Bacillus subtilis]	78	64	069
995 1 2	978	727	gi 296947	uridine kinase [Escherichia coli]	78	64	252
1045 1	e -	1 401	gi 1407784	orf-1; novel antigen [Staphylococcus aureus]	78	61	399
1163 2	368	186	gi 410117	diaminopimelate decarboxylase [Bacillus subtilis]	78	54	183
2191 1	1 794	399	gi 215098	excisionase [Bacteriophage 154a]	78	65	396
2933 1	7	181	gi 1204436	pyruvate formate-lyase [Haemophilus influenzae]	78	73	180
3041 2	129	1 317	gi 624632	GltL [Escherichia coli]	78	53	189
3581 1	105	401	gi 763186	3-ketoacy1-coA thiolase [Saccharomyces cerevisiae]	78	55	297
3709 1	8	1 230	gi 460689	TVG [Thermoactinomyces vulgaris]	78	58	228
3974 1	1 528	1 265	gi 558839	unknown [Bacillus subtilis]	78	65	264
3980 1	e 	1 401	gi 39956	IIGIc (Bacillus subtilis)	78	62	399
4056 1	647	1 354	gi 1256635	dihydroxy-acid dehydratase [Bacillus subtilis]	78	55	294
4114 1	1 630	316	pir 809372 8093	hypothetical protein - Trypanosoma brucei	78	62	315
4185 1		179	lgi 1339950	large subunit of NADH-dependent glutamate synthase [Plectonema boryanum]	78	58	177

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	+ uis *	* ident	length (nt)
4235	-	655	1 329	gi 558839	unknown [Bacillus subtilis]	78	1 09	327
4352		541	302	gi 603768 	Hutl protein, imidazolone-5-propionate hydrolase [Bacillus subtilis] gi 603768 Hutl protein, imidazolone-5-propionate hydrolase Bacillus subtilis]	78	63 -	240
4368		612	1 307	gi 1353678	heavy-metal transporting P-type ATPase [Proteus mirabilis]	78 1	59	306
4461		1 428	216	gi 1276841	glutamate synthase (GOGAT) [Porphyra purpurea]	78	36	213
1 4530		474	1 238	gi 39956		78 1	65	237
3	7	1 2969	1 2073	gi 1109684	ProV [Bacillus subtilis]	7.7	56	1 268
1 12	7	1 2426	1 1965	gi 467335	ribosomal protein L9 (Bacillus subtilis)	1 11	59	462
1 27		2	1 388	gi 1212728	YqhI [Bacillus subtilis]	1 77	63	387
39	7	290	1252	gi 40054	phenylalanyl-tRNA synthetase beta subunit (AA 1-804) (Bacillus btilis)	1 77	1 09	663
42	·	2704	1 2931		30s ribosomal subunit protein S14 [Escherichia coli] sp P02370 RS14_ECOLI 30s RIBOSOWAL PROTEIN S14. (SUB 2-101)	1.77	65	228
46	18	115459	116622	gi 297798 	mitochondrial formate dehydrogenase precursor [Solanum tuberosum] pir JQ2272 JQ2272 formate dehydrogenase (EC 1.2.1.2) precursor, itochondrial - potato	1	55	1164
100	4	1 4562	1 4002	gi 1340128	ORF1 [Staphylococcus aureus]	177	54	561
102	8	5378	5713	gi 1311482	acetolactate synthase [Thermus aquaticus]	1 77	57	336
109	7	4742	5383	gi 710637	Unknown (Bacillus subtilis)	77	56	642
117	1	5	1228	gi 1237015	ORF4 [Bacillus subtilis]	1 77	53	1227
124	110	8323	1 7688	gi 405819	[thymidine kinase [Bacillus subtilis]	177	63	636
147	m -	1146	1 985	gi 849027	hypothetical 15.9-kDa protein [Bacillus subtilis]	1 77	37	162
152	. 10	7354	1 7953	gi 1205583 	spermidine/putrescine transport ATP-binding protein [Haemophilus influenzae]	77	22	009
169	7	1004	1282	gi 473825	'elongation factor EF-Ts' [Escherichia coli]	77	58	279
1 184	- 5	380	11147	gi 216314		1 17	09	768
189	7	3296	1 3868	gi 853809	ORF3 Clostridium perfringens	77	48	573
193		132	1 290	gi 1303788	YqeH [Bacillus subtilis]	1 77	54	159
195	æ	8740	8414	gi 1499620	[M. jannaschii predicted coding region MJ0798 [Methanococcus jannaschii]	1	44	327
1 205	8	1 5428	5204	gi 216340	ORF for adenylate kinase (Bacillus subtilis)	77	61	225

Table?

S. aureus - Putative coding regions of novel proteins similar to known proteins

		-		-				
Contig	g ORF	Start (nt)	Stop (nt)	match acession	match gene name	8 sim	% ident	length (nt)
1 205	129	114795	114502	gi 786155	Ribosomal Protein L23 (Bacillus subtilis)	177	62	294
211		1 1908	1 2084	gi 410132		17	47	177
1 217	ر ا	3478	4416	gi 496254	fibronectin/fibrinogen-binding protein [Streptococcus pyogenes]	7.2	54	686
1 232		1 267	866	gi 1407784	orf-1; novel antigen (Staphylococcus aureus)	7.7	57	732
233	7	1 1819	1346	gi 467408	unknown (Bacillus subtilis)	77	61	474
243	e -	1 2661	1 2299	 gi 516155	unconventional myosin [Sus scrofa]	1 77	32	363
299		89	692	gi 467436	unknown [Bacillus subtilis]	77	54	702
301	4	1468	1283	gi 950071	HTP-bind. pyrimidine kinase [Mycoplasma capricolum] pir S48605 S48605 hypothetical protein - Mycoplasma capricolum SGC3) (fragment)	1 77	48	186
302	2	1 2741	3211	gi 508980	pheB [Bacillus subtilis]	1 11	57	471
302	7	3832	1 4863	gi 147783	ruvB protein [Escherichia coli]	77	09	1029
307	6	5402	4797	gi 1070015		7.7	09	909
312		66	1391	l -d	malic enzyme (EC 1.1.1.38) [Bacillus stearothermophilus] pir[A33307 DEBSXS malate dehydrogenase oxaloacetate-decarboxylating) (EC 1.1.1.38) - Bacillus tearothermophilus	77	62	1293
312		1541	1 2443	gi 1399855		77	28	1 606
321	- 5	1 5666	1 4596	gi 39844	fumarase (citG) (aa 1-462) [Bacillus subtilis]	7.7	65	1071
354		47	1 568	gi 1154634	YmaB [Bacillus subtilis]	77	57	522
365		7	1021	gi 143374 	phosphoribosyl glycinamide synthetase (PUR-D; gtg start codon) Bacillus subtilis]	77	62	1020
374			1 708	gi 1405446	transketolase [Bacillus subtilis]	17	61	708
385	٦.	1128	1 565	gi 533099	 endonuclease III [Bacillus subtilis]	177	63	564
392		594	1940	gi 556014 	UDP-N-acetyl muramate-alanine ligase [Bacillus subtilis] sp P40778 MURC_BACSU UDP-N-ACETYLMURAMATEALANINE LIGASE (EC .3.2.8) (UDP-N- ACETYLMURANOYL-L-ALANINE SYNTHETASE) (FRAGMENT):	77	65	1347
405	5	1 4079	1 3570	gi 1303912	YqhW [Bacillus subtilis]	77	64	510
487	4	1302	1472	gi 432427	ORF1 gene product [Acinetobacter calcoaceticus]	17	48	171
522		5	1 562	pir A01179 SYBS	ir A01179 SYBS tyrosinetRWA ligase (EC 6.1.1.1) - Bacillus stearothermophilus	77	63	561

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF IID	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
523	~	1587	1351	gi 1387979	144% identity over 302 residues with hypothetical protein from Synechocystis l sp, accession D64006 CD; expression induced by environmental stress; some similarity to glycosyl transferases; two potential membrane-spanning helices [Bacillus subtil	1 77	4 1 80 1 00	237
536	7	983	612	gi 143366 	adenylosuccinate lyase (PUR-B) [Bacillus subtilis] pir C29326 WZBSDS adenylosuccinate lyase (EC 4.3.2.2) - Bacillus ubtilis	+ 77	61	372
548	7	339	1 872	gi 143387	aspartate transcarbamylase [Bacillus subtilis]	1 77	56	534
1 597	7	1 2	481	gi 904198	hypothetical protein (Bacillus subtilis)	1 77	33 +	480
1 633	1 2	1747	1313	gi 387577	ORFIA [Bacillus subtilis]	1	64	435
1 642	1	85	1 360	gi 46971	epiP gene product [Staphylococcus epidermidis]	77	61	276
629	7	125	1219	gi 1072381	glutamyl-aminopeptidase [Lactococcus lactis]	1 77	62	1095
0.29	4	1587	1 1820	gi 1122760	unknown [Bacillus subtilis]	1. 11	58	234
189		7	1 391	gi 1377823	aminopeptidase [Bacillus subtilis]	1 11	65	390
815		10	1 573	gi 1303861	YqgN [Bacillus subtilis]	1 11	49	564
668	-		1 225	gi 1204844	H. influenzae predicted coding region HIO594 [Haemophilus influenzae]	1 77	55	225
1 1083	-	8	188	gi 460828	18969 [Saccharomyces cerevisiae]	1 77	99	186
1942		415	209	gi 160047 	p101/acidic basic repeat antigen [Plasmodium falciparum] pir A29232 A29232 101K malaria antigen precursor - Plasmodium alciparum (strain Camp)	77,	38	207
1 2559	1		171	gi 1499034	IM. jannaschii predicted coding region MJ0255 [Methanococcus jannaschii]	1 77	61	171
2933		243	401	gi 42370 	Ipyruvate formate-lyase (AA 1-760) [Escherichia coli] ir S01788 S01788 formate C-acetyltransferase (EC 2.3.1.54) - cherichia coli	1 77	72	159
1 2966		56	1 292	gi 1524397	glycine betaine transporter OpuD [Bacillus subtilis]	17	45	237
2976		614	309	gi 40003 	oxoglutarate dehydrogenase (NADP+) [Bacillus subtilis] p P23129 0D01 BACSU	1 77	09	306
1 2979	7	678	1 400	gi 1204354	spore germination and vegetative growth protein [Haemophilus influenzae]	1.77	61	279
1 2988		601	1 377	 gi 438465 	Probable operon with orff. Possible alternative initiation codon, ases 2151-2153. Homology with acetyltransferases.; putative Bacillus subtilis]	77	35	225
1 2990		331	167	gi 142562 	ATP synthase epsilon subunit [Bacillus megaterium] pir B28599 PWBSEM H+- transporting ATP synthase (EC 3.6.1.34) psilon chain - Bacillus megaterium	77	63	165
3032	. –	3	1 389	gi 488430	alcohol dehydrogenase 2 [Entamoeba histolytica]	1 77	56	387
1 3057			195	gi 468764	mocR gene product [Rhizobium meliloti]	1 77	50	195
•	-			_	_			

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig ORF	ORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	* ident	length (nt)
4008		726	400	gi 603768	HutI protein, imidazolone-5-propionate hydrolase [Bacillus subtilis] [gi[603768 HutI protein, imidazolone-5-propionate hydrolase Bacillus subtilis]	1 66	52	327
4048	1	703	386	gi 216278	gramicidin S synthetase 1 [Bacillus brevis]	7.7	55	318
4110	- 1	3	1 368	pir S52915 S529	Initrate reductase alpha chain - Bacillus subtilis (fragment)	177	61	366
4115	1		348	lgi1517205	167 kDa Myosin-crossreactive streptococcal antigen [Streptococcus yogenes]	1 11	1 59	348
4225		290	297	gi 1322245	mevalonate pyrophosphate decarboxylase [Rattus norvegicus]	1 11	1 09	294
1 4611	- 5	494	1 327	gi 508979	GTP-binding protein [Bacillus subtilis]	17	57	168
4668	-	1 361	1 182	pir S52915 S529	Initrate reductase alpha chain - Bacillus subtilis (fragment)	1 77	61	180
25	1	1 2	1627	gi 1150620	MmsA [Streptococcus pneumoniae]	76	28	1626
38	2	1488	1 2537	pir A43577 A435	regulatory protein pfoR - Clostridium perfringens	76	57	1050
1 52	5	1 2962	4041	gi 1161061	dioxygenase [Methylobacterium extorquens]	76	62	1080
1 56	120	127389	127955	gi 467402	unknown [Bacillus subtilis]	76	95	1 295
57	115	112046	112219	gi 1206040	weak similarity to keratin [Caenorhabditis elegans]	192	40	174
91	7	1 1062	1 2261	gi 475715	acetyl coenzyme A acetyltransferase (thiolase) [Clostridium cetobutylicum]	1 9/	57	1200
86	1 2	818	1 1624	gi 467422	unknown [Bacillus subtilis]	76	62	807
86	- 5	1 2965	1 3228	gi 897793	1998 gene product [Pediococcus acidilactici]	192	52	264
86	80	1 5922	6326	gi 467427	methionyl-tRNA synthetase [Bacillus subtilis]	76	53	405
104	m 	1322	1885	gi 216151 	DNA polymerase (gene L; ttg start codon) [Bacteriophage SP02] gil579197 SP02 DNA polymerase (aa 1-648) [Bacteriophage SP02] pir A21498 DJBPS2 DNA- directed DNA polymerase (EC 2.7.7.7) - phage P02	1 94	63	564
124	6	8134	7055	gi 853776 	peptide chain release factor 1 [Bacillus subtilis] pir S55437 S55437 peptide chain release factor 1 - Bacillus ubtilis	76	28	1080
164	- 5	2832	3311	gi 1204976	prolyl-tRNA synthetase [Haemophilus influenzae]	16	53	480
168	1 2	1 2617	1841	gi 1177253		76	58	1777
1 189	1 2	163	888	gi 467384	unknown [Bacillus subtilis]	76	63	726
235	e – –	2253	3518	gi 142936 	folyl-polyglutamate synthetase [Bacillus subtilis] pir B40646 B40646 folC - Bacillus subtilis	76	53	1266
236		1 335	925	gi 1146197	putative [Bacillus subtilis]	192	54	591
1 237	80	1 5323	5541	gi 1279261	[F13G3.6 [Caenorhabditis elegans]	76	47	219
	1							

Table 2

1035 1293 1236 471 1503 624 264 906 744 222 528 306 723 498 720 258 735 225 183 579 243 180 | % sim | % ident | length (nt) 49 72 61 09 22 57 36 44 52 47 55 28 29 65 9 29 28 61 61 26 38 25 91 91 9/ 9/ 16 16 91 91 97 16 97 91 16 91 91 16 putative membrane spanning subunit [Bacillus subtilis] pir|552382|552382 | probable membrane spanning protein - Bacillus ubtilis dihydrolipoamide transsuccinylase (odhB; EC 2.3.1.61) [Bacillus ubtilis] M. jannaschii predicted coding region MJ1322 [Methanococcus jannaschii] streptococcal pyrogenic exotoxin type C (speC) precursor Streptococcus [39.0% identity to the Escherichia coli S1 ribosomal protein; putative adenylosuccinate synthetase (Bacillus subtilis) splP29726|PURA BACSU ADENYLOSUCCINATE SYNTHETASE (EC 6.3.4.4) IMP--ASPARTATE LIGASE). 6-phospho-1-fructokinase (gtg start codon; EC 2.7.1.11) [Bacillus S. aureus - Putative coding regions of novel proteins similar to known proteins |alternate gene name mgt; CG Site No. 497 [Escherichia coli] | pir|S56468|S56468 mgtA protein - Escherichia coli |dihydrodipicolinate synthase [Methanococcus jannaschii] [No definition line found [Mycobacterium leprae] |acetylornithine deacetylase [Escherichia coli] |major capsid protein [Human cytomegalovirus] |transport protein [Salmonella typhimurium] |hypothetical protein [Bacillus subtilis] |ORF YKL027w [Saccharomyces cerevisiae] |spoOF gene product [Bacillus subtilis] OppD homologue [Rhizobium sp.] |CapJ [Staphylococcus aureus] (ClpB [Synechococcus sp.] [AlsT [Bacillus subtilis] |MutS [Bacillus subtilis] {YqfG [Bacillus subtilis] |YqkG [Bacillus subtilis] yeeE [Escherichia coli] [Bacillus subtilis] | tearothermophilus] | match gene name pyogenes] acession | 4585 |gi|1510348 |gi|1146215 1gi11303823 |gi|1002520 |gi|1486422 |gi|1511333 |gi|1303984 |gi|1405464 match | 1794 |gi|666982 | 4624 |gi|143312 1gi1405956 |gi|396304 |gi|537084 | 4 | 4331 | 4858 |gi|143268 |gi|486025 |gi|153821 gi1885934 |gi|330613 |gi|904199 |gi|142443 |gi|467154 |gi|506706 |gi|153898 |gi|40177 1701 2854 1036 1907 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 4492 3302 1 307 1 798 1 290 1 206 1 266 1 557 1 228 580 222 995 735 1 244 5490 3611 1051 1 4022 4356 1054 2568 1158 16 1 1 409 1 1 1 273 12 | 175 1 547 452 24 1563 | 1 | 529 1 1 486 1 1 672 7 _ 11 1 2 4 --7 - 2 -- 2 ~ 4 _ 404 472 602 619 997 263 304 347 358 411 591 999 962

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

+					+	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	++	+
Contig	g lORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	8 ident	length (nt)
1 2942		1 29	400	gi 710020	nitrite reductase (nirB) [Bacillus subtilis]	16	1 65	372
1 2957	1	1 377	216	gi 1511251	hypothetical protein (SP:P42404) [Methanococcus jannaschii]	76	47	162
1 2980		1 554	279	gi 1405464	AlsT [Bacillus subtilis]	76	53	276
3015		1 649	326	gi 408115	ornithine acetyltransferase [Bacillus subtilis]	76	61	324
3124		13	174	gi 882705	ORF_0401 [Escherichia coli]	76	65	162
3179		m 	161	gi 168477	ferredoxin-dependent glutamate synthase [Zea mays] pir A38596 A38596 glutamate synthase (ferredoxin) (EC 1.4.7.1) - aize	91	53	159
3789		2	379	gil39956	IIGlc (Bacillus subtilis)	76	1 25 1	378
3892		8 -	314	gi 1510398	[ferripyochelin binding protein [Methanococcus jannaschii]	76	52	312
1 3928		1 798	1 400	gi 143016		9/	65	399
4159		1 757	1 386	sp P80544 MRSP_	METHICILLIN-RESISTANT SURFACE PROTEIN (FRAGMENTS).	9/	99	372
1 4204		1 17	331	gi 296464	ATPase [Lactococcus lactis]	76	56 1	315
1 4398		1 494	1 249	gi1987255	Menkes disease gene [Homo sapiens]	97	48	246
4506		- 2	313	gi 216746	D-lactate dehydrogenase [Lactobacillus plantarum]	92	4 4 4	312
4546		477	1 247	gi 1339950	large subunit of NADH-dependent glutamate synthase [Plectonema boryanum]	76	61	231
4596		1 379	191	1911560027	cellulose synthase [Acetobacter xylinum]	76	1 0/	189
4	5	1 5257	1 4337	gi 882532	ORF_0294 [Escherichia coli]	7.5	59	921
9		164	952	gi 40960	OTCase [Escherichia coli]	75	56	789
12	е —	1 5935	1 3944	gi 467336	unknown [Bacillus subtilis]	75	57	1992
1 23	118	118272	117310	gi 1296433	O-acetylserine sulfhydrylase B (Alcaligenes eutrophus)	7.5	55	963
1 25	e 	1 2356	1 3393	gi 1502419		7.5	1 26 1	1038
36	8	1 5765	1 6037	gi 1256517	unknown [Schizosaccharomyces pombe]	75	45	273
1 46	113	111186	112058	gi 48972	Initrate transporter [Synechococcus sp.]	7.5	1 46 1	873
- 51	7 + -	1 3474	13677	gi 143607	sporulation protein (Bacillus subtilis)	75	61	204
53	116	116850	116590	gi 143402 	recombination protein (ttg start codon) [Bacillus subtilis] gi 1303923 RecN [Bacillus subtilis]	75	51	261
1 74	e 	3572	1 2568	gi 1204847	ornithine carbamoyltransferase [Haemophilus influenzae]	75	61	1005
1 1 1 1 1 1 1	1	 	1			 	+	

| % sim | % ident | length (nt) 54 48 54 55 99 59 63 53 61 51 22 75 22 62 52 28 9 53 58 49 23 54 75 75 75 75 75 75 75 75 75 75 75 75 154.8% identity with Neisseria gonorrhoeae regulatory protein PilB; putative | phosphoribosyl aminoidazole succinocarboxamide synthetase (PUR-C; tg start Ibranched chain aa transport system II carrier protein [Haemophilus S. aureus - Putative coding regions of novel proteins similar to known proteins |HYPOTHETICAL PROTEIN IN PURB 5'REGION (ORF-15) (FRAGMENT) |putative ppGpp synthetase [Streptomyces coelicolor] |ORF-2 upstream of gbsAB operon [Bacillus subtilis] [homologous to sp:PHOR_BACSU [Bacillus subtilis] |porphobilinogen deaminase [Bacillus subtilis] Initrite extrusion protein (Bacillus subtilis) [ATP-dependent nuclease [Bacillus subtilis] |seryl-tRNA synthetase [Bacillus subtilis] [[gap] gene products [Bacillus megaterium] | Ihypothetical protein [Synechocystis sp.] |hypothetical protein [Synechocystis sp.] [ATP-binding protein [Bacillus subtilis] (clwD gene product [Bacillus subtilis] |function unknown [Bacillus subtilis] [unknown [Schizosaccharomyces pombe] | codon) [Bacillus subtilis] |unknown (Bacillus subtilis) [unknown [Bacillus subtilis] |YqiA (Bacillus subtilis] (NrdE [Bacillus subtilis] [YbbF [Bacillus subtilis] |MutL [Bacillus subtilis] [Bacillus subtilis] I match gene name | sp | P25745| YCFB acession 6625 | 7530 |gi|1303916 |gi|1064813 |gi|1064810 |gi|1001824 |gi|1147593 1gi11177251 |gi|1524394 |gi|1001195 |gi|1256135 gi | 1052800 |gi|1205958 |gi|1154632 |gi|1256654 |gi|1002521 |gi|142440 | 3930 |gi|143368 |gi|143367 |gi|405622 |gi|143316 | 3185 |gi|467403 |gi|755153 |gi|143037 | 1714 |gi|971337 | 3 | 1314 | 1859 | gil467405 1139 1 1 | 2240 | 1122 1 4878 1 2340 | 3590 110649 1 5903 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 6084 | 6896 5252 4221 1 5798 2995 1107 9374 1 5620 1844 | 1503 143 4596 389 847 4628 5588 1 4480 2024 9529 2605 0069 1748 1 4353 114 111266 1 2050 18 1 5760 5439 3819 283 81 1 518 159 <u>~</u> 2 œ — <u>~</u> 9 ~ m _ _ 5 7 110 4 _ 10 4 7 110 130 82 85 85 87 87 108 120 121 124 128 161 172 195 200 206

1296

360 825 918

156

1992 1281 231 819 1197 1119

1251

813 342

711

669

1980

900 618 2172

1059

144

141

Table 2

| % ident | length (nt) | & sim |methyl coenzyme M reductase system, component A2 [Methanococcus jannaschii] | pir|JC4110|JC41 |triacylglycerol lipase (EC 3.1.1.3) 2 - Mycoplasma mycoides subsp. mycoides 44% identity over 302 residues with hypothetical protein from Synechocystis sp, accession D64006_CD; expression induced by environmental stress; some similarity to glycosyl transferases; two potential membrane-spanning helices [Bacillus subtil Matches PS00017: ATP GTP A and PS00301: EFACTOR GTP; similar to longation M. jannaschii predicted coding region MJ1651 [Methanococcus jannaschii] | factor G, TetM/TetO tetracycline-resistance proteins Escherichia coli] S. aureus - Putative coding regions of novel proteins similar to known proteins undefined open reading frame [Bacillus stearothermophilus] |phoP protein (put.); putative [Bacillus subtilis] lepiP gene product [Staphylococcus epidermidis] |acetolactate synthase [Klebsiella pneumoniae] |potF gene product [Clostridium perfringens] |tegument protein [Saimiriine herpesvirus 2] |ribonuclease P [Saccharomyces cerevisiae] |Pyruvate Kinase [Bacillus psychrophilus] |OMP-PRPP transferase (Bacillus subtilis) citrate synthase II (Bacillus subtilis) OMP decarboxylase [Lactococcus lactis] |dihydroorotase [Bacillus caldolyticus] |spoIIIEA protein [Bacillus subtilis] |surfactin [Haemophilus influenzae] |EpiG [Staphylococcus epidermidis] |unknown [Pseudomonas aeruginosa] |spec [Streptococcus pyogenes] |ORF_0236 [Escherichia coli] |YqhI [Bacillus subtilis] match gene name acession 1gi | 1511246 |gi|1511604 1gi11205328 |gi|1183839 |gi|1387979 |gi|1176401 |gi|1041097 gi | 1212728 |gi|1070361 |gi|1149682 |gi|143394 |gi|285621 |gi|487433 1gi1304976 |gi|312441 |gi|143582 1gi1143328 |gi|529754 |gi|882541 |gi|149211 gi1330993 |gi|172442 |gi|46971 1 3387 1 2088 (nt) 1 164 1 832 Contig |ORF | Start | ID | ID | (nt) | 1 1 1 1662 1 2437 1 4321 1 353 1 127 1 430 1 494 m m <u>۳</u> - 5 -~ --_ S ~

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	1 ORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
754		7	481	gi 1303901	YqhT (Bacillus subtilis)	75	57	480
1 763	7	1 563	1 393	gi 1205145	multidrug resistance protein (Haemophilus influenzae)	75	51	171
175		961	1 482	pir B36889 B368	leuA protein, inactive - Lactococcus lactis subsp. lactis (strain IL1403)	75	63	480
1 793			1 180	gi 143316	[gap] gene products [Bacillus megaterium]	75	57	180
008	-	318	160	gi 509411	NFRA protein [Azorhizobium caulinodans]	75	34	159
811		1117	1 560	gi 143434	Rho Factor [Bacillus subtilis]	75	09	558
1 940	-	1 493	1 329	gi 1276985	arginase [Bacillus caldovelox]	75	50 1	165
971	7	1 37	1 252	gi 1001373	hypothetical protein [Synechocystis sp.]	75	58	216
1059		384	1 232	gi 726480	L-glutamine-D-fructose-6-phosphate amidotransferase [Bacillus ubtilis]	75	67	153
1109	~	219	374	gi 143331 	alkaline phosphatase regulatory protein (Bacillus subtilis) pir A27650 A27650 regulatory protein phoR - Bacillus subtilis sp P23545 PHOR_BACSU ALKALINE PHOSPHATASE SYNTHESIS SENSOR PROTEIN HOR (EC 2.7.3).	75	23	156 -
1268		271	137	gi 304135	ornithine acetyltransferase (Bacillus stearothermophilus) spiQ07908 ARGJ BACST GLUTAMATE N-ACETYLTRANSFERASE (EC 2.3.1.35) ORNITHINE ACETYLTRANSFERASE) (ORNITHINE TRANSACETYLASE) / MINO-ACID ACETYLTRANSFERASE (EC 2.3.1.1) (N-ACETYLGLUTAMATE YNTHA	75	63	135
1500	-	324	163	gi 1205488	excinuclease ABC subunit B [Haemophilus influenzae]	75	57	162
1529		1 798	1 400	gi 1002521	Mutl [Bacillus subtilis]	75	54	399
3010		1 770	1 387	gi 1204435	pyruvate formate-lyase activating enzyme [Haemophilus influenzae]	75	54	384
3105			180	gi 1041097	Pyruvate Kinase [Bacillus psychrophilus]	75	57	180
3117		45	212	gi 899317 	peptide synthetase module [Microcystis aeruginosa] pir[549111]549111 probable amino acid activating domain - icrocystis aeruginosa (fragment) [SUB 144-528]	75	42	168
3139	- 5	139	345	gi 145294	adenine phosphoribosyl-transferase [Escherichia coli]	75	1 99	207
3880		1 618	310	gi 1009366	Respiratory nitrate reductase [Bacillus subtilis]	75	58	309
3911		- 48	401	gi 433991	ATP synthase subunit beta [Bacillus subtilis]	75	89	354
3957		2	379	pir D36889 D368	13-isopropylmalate dehydratase (EC 4.2.1.33) chain leuc - Lactococcus lactis subsp. lactis (strain IL1403)	75	65	378
1 4005		. —	1 259	gi 216746	D-lactate dehydrogenase [Lactobacillus plantarum]	75	48	255
1 4080		1 73	1 333	gi 415855	deoxyribose aldolase [Mycoplasma hominis]	75	59	261
-							-	

Table 2

339 300 333 291 246 357 309 069 1452 645 216 919 1299 1482 555 1596 276 351 564 972 462 621 1212 705 | % ident | length (nt) 57 26 49 22 52 55 55 57 47 52 09 54 89 20 57 29 57 49 28 21 57 28 45 74 75 75 75 75 74 74 74 74 74 74 74 74 74 8 Sim |ATP binding protein of transport ATPases [Bacillus firmus] ir|S15486|S15486 | ATP-binding protein - Bacillus firmus p|P26946|YATR_BACFI HYPOTHETICAL | ATP-BINDING TRANSPORT PROTEIN. |hsdM gene of EcoprrI gene product [Escherichia coli] pir|538437|538437|hsdM | protein - Escherichia coli pir|S09629|S09629 hypothetical protein A -|methylated-DNA--protein-cysteine methyltransferase [Haemophilus influenzae] ribonucleoside triphosphate reductase [Escherichia coli] pir|A47331|A47331 |large subunit of NADH-dependent glutamate synthase [Plectonema boryanum] |dipeptide transport system permease protein [Haemophilus influenzae] anaerobic ribonucleotide reductase - Escherichia oli [major surface glycoprotein [Pneumocystis carinii] |phosphoglycerate kinase [Bacillus megaterium] |endoglucanase [Methanococcus jannaschii] |hypothetical protein [Bacillus subtilis] |aspartate-tRNA ligase (Escherichia coli) Initrate transporter [Synechococcus sp.] [DNA-binding protein [Bacillus subtilis] |devA gene product [Anabaena sp.] Escherichia coli (SUB 40-520) (ttg start [Campylobacter coli) [putative [Lactococcus lactis] [putative [Lactococcus lactis] [putative [Lactococcus lactis] |histidase [Bacillus subtilis] [ATPase [Lactococcus lactis] |unknown [Bacillus subtilis] |YqhJ [Bacillus subtilis] |TreP (Bacillus subtilis] [NupC [Escherichia coli] acession |gi|1339950 |gi|1239988 |gi|1205430 |gi|1107528 |gi|1256653 1gi11377833 |gi|1000451 |gi|1212729 |gi|450688 |gi|1204652 |gi|149435 |gi|1510631 |gi|143318 |gi|294260 gi | 296464 1gi | 443793 |gi|149429 1gi|149435 |gi|146970 1gi | 143076 |gi|407773 1gi148972 |gi|41015 1gi | 39478 114419 1368 6126 6221 4848 114897 13136 3902 8574 | 7093 1 4803 6581 5791 339 112221 1 1622 1 5150 Contig | ORF | Start | Stop (nt) 374 336 359 1 312 1 277 318 113121 (nt) 5776 110770 2266 114334 1 5961 2320 5237 6910 5063 12561 4 | 4873 7 | 4342 6745 602 664 73 1 1 1 715 1 620 664 --1 1 | 552 -. -- 2 œ <u>ه</u> 7 9 18 115 116 -5 7 ᠬ _ <u>~</u> 7 4111 4136 4144 4237 4306 4343 4552 유 38 20 26 64 99 67 102 102 120 121 128 130 136 145 152

Table ?

o

1137 312 453 1008 561 1188 507 1239 309 1050 555 612 1065 747 432 168 1191 234 255 1002 942 333 | % ident | length (nt) 46 55 22 64 42 54 53 57 26 65 51 99 25 28 52 64 63 62 74 l % sim pir|E39845|E39845 carbamoyl-phosphate synthase glutamine-hydrolyzing) (EC 6.3.5.5), pyrimidine-repressible, small hain - Bacillus subtilis |threonine synthase (thrC) (AA 1-352) [Bacillus subtilis] ir|A25364|A25364 |glutamate-1-semialdehyde 2,1-aminotransferase [Bacillus subtilis]
| pir[D42728|D42728 glutamate-1-semialdehyde 2,1-aminomutase (EC .4.3.8)
| Bacillus subtilis glutaminase of carbamyl phosphate synthetase [Bacillus subtilis] hypothetical protein (GB:U14003_302) [Haemophilus influenzae] |hypothetical protein (SP:P25745) [Haemophilus influenzae] single strand DNA binding protein [Bacillus subtilis] |glycine betaine transporter OpuD [Bacillus subtilis] threonine synthase (EC 4.2.99.2) - Bacillus btilis |Na/H antiporter system ORF2 [Bacillus alcalophilus] |glycyl-tRNA synthetase [Methanococcus jannaschii] lalpha-acetolactate synthase [Lactococcus lactis] |malate thiokinase [Methylobacterium extorquens] (ATPase subunit a [Bacillus stearothermophilus] [ATP-binding protein [Haemophilus influenzae] [homoserine dehydrogenase [Bacillus subtilis] [acid glycoprotein [Streptococcus pyogenes] lipa-58r gene product [Bacillus subtilis] |ribosomal protein L6 [Bacillus subtilis] |ORF3 gene product [Bacillus subtilis] |CbrB protein [Erwinia chrysanthemi] quinol oxidase (Bacillus subtilis) (thioredoxin [Bacillus subtilis] |unknown [Rhizobium meliloti] [putative [Bacillus subtilis] |YqhZ [Bacillus subtilis] acession |gi|1204430 |gi|1044979 |gi|1204872 |gi|1205579 |gi|1303915 1gi11499005 |gi|1524397 |gi|1146207 |gi|142520 |gi|217040 |gi|143040 1gi1854656 |gi|467374 1gi1809542 |gi|143398 |gi|143389 | 1128 |gi|534857 1gi1694121 |gi|473902 |gi|413982 |gi|558494 |gi|40211 1 4665 1311 111064 | 111375 5612 110339 6506 6710 3334 1 4081 1 3270 11193 1740 1 2799 5313 1 1407 (nt) 999 | 1 862 942 734 320 5574 5714 (nt) 3109 6064 6551 1 2503 111346 9619 4521 1 4389 1220 1473 1742 1 4083 3305 11174 382 574 1 739 402 ო Contig | ORF | 110 6 111 117 <u>س</u> 9 _ 9 - 1 7 7 <u>۔</u> د 7 _ ~ 164 175 195 205 241 249 261 278 309 405 433 452 461 461 462 478 501

Table 2

219 1167 327 156 189 255 186 177 303 168 477 483 186 258 282 492 366 237 399 288 222 54 59 45 55 20 47 46 54 48 36 09 28 26 51 64 41 58 55 62 28 62 57 74 74 l % sim | pir|D42728|D42728 glutamate-1-semialdehyde 2,1-aminomutase (EC .4.3.8) | Bacillus subtilis |phosphoribosyl glycinamide synthetase (PUR-D; gtg start codon) Bacillus |M. jannaschii predicted coding region MJ0938 [Methanococcus jannaschii] 121.4% of identity to trans-acting transcription factor of Sacharomyces cerevisiae; 25% of identity to sucrose synthase of Zea mays; putative glutamate-1-semialdehyde 2,1-aminotransferase [Bacillus subtilis] |pir|552915|5529 |nitrate reductase alpha chain - Bacillus subtilis (fragment) 13-phosphoglycerate kinase [Thermotoga maritima] epiD gene product [Staphylococcus epidermidis] |peptide-synthetase ORF1 [Bacillus subtilis] |sbcC gene product [Clostridium perfringens] |ORF1; putative [Streptococcus parasanguis] Ihypothetical protein [Synechocystis sp.] lipa-24d gene product (Bacillus subtilis) |recJ gene product [Erwinia chrysanthemi] |unknown [Staphylococcus haemolyticus] [ttg start [Campylobacter coli] [aconitase [Bacillus subtilis] |ORF3 [Lactococcus lactis] |ORF V (Bacillus subtilis) |orf6 [Lactobacillus sake] |YqkM [Bacillus subtilis] [YqhL [Bacillus subtilis] orf4 [Bacillus subtilis] [YqgF [Bacillus subtilis] [Bacillus subtilis] | subtilis acession |gi|1006605 |gi|1303853 |gi|1146237 |gi|1072380 gi|1303893 |gi|1107528 gil1510977 |gi|1370207 gi|1303994 gi|1149701 gi|1405454 gi | 143040 gi | 520844 |gi|580910 |gi|143374 |gi|450686 |gi|153833 |gi|143458 |gi|413948 |gi|495245 gi|46970 12 | 1269 | 1595 1298 11758 188 1 256 189 1 231 1 477 259 724 360 492 528 452 238 400 363 191 305 304 221 Contig |ORF | Start | ID | ID | (nt) | 1 2924 1780 373 817 1 650 1 1 | 510 1 1 1 374 1 471 373 443 184 607 4587 | 1 | 458 604 1 798 - _ --7 7 -7 7 1106 1135 1234 2586 2959 2983 4035 4045 4283 2962 3062 4449 686 573 836 296 724

 | % ident | length (nt) | % sim [alpha-isopropylmalate isomerase (put.); putative [Rhizomucor ircinelloides] ribonucleoside triphosphate reductase [Escherichia coli] pir|A47331|A47331 - acillus |anaerobic ribonucleoside-triphosphate reductase [Haemophilus influenzae] IM. jannaschii predicted coding region MJ0775 [Methanococcus jannaschii] glutamate synthase large subunit (EC 2.6.1.53) [Escherichia coli] | pir|A29617|A29617 glutamate synthase (NADPH) (EC 1.4.1.13) large hain phosphoribosyl aminoimidazole carboxy formyl ormyltransferase/inosine |meso-2,3-butanediol dehydrogenase (D-acetoin forming) [Klebsiella 13-isopropylmalate dehydrogenase (AA 1 - 365) [Bacillus subtilis] pir|A26522|A26522|A26522 | 1.1.1.85) S. aureus - Putative coding regions of novel proteins similar to known proteins monophosphate cyclohydrolase (PUR-H(J)) Bacillus subtilis) threonine dehydratase 2 (EC 4.2.1.16) [Escherichia coli] | anaerobic ribonucleotide reductase - Escherichia oli |dihydroorotate dehydrogenase [Agrocybe aegerita] ornithine cyclodeaminase [Rhizobium meliloti] lipa-76d gene product [Bacillus subtilis] |stage V sporulation [Bacillus subtilis] lipa-7d gene product [Bacillus subtilis] |murD gene product [Bacillus subtilis] [pepT gene product [Bacillus subtilis] [moaB gene product [Escherichia coli] |putative [Bacillus subtilis] |unknown [Bacillus subtilis] |Prow [Bacillus subtilis] [YbbF [Bacillus subtilis] AppF [Bacillus subtilis] |yeiN [Escherichia coli] Escherichia coli match gene name pneumoniae) acession |gi|1256135 |gi|1468939 |gi|1109685 | 3493 | 2600 | gi|1510849 | 5726 | 6223 |gi|1204333 |gi|1429259 match |gi|146970 |gi|146208 1gi1143727 1gi1297060 |gi|467442 | 4914 | | 414000 lgi1405885 |gi|580895 |gi|147923 1gi1677944 |gi|580932 |gi|580891 |gi|166338 |gi|143373 |gi|168367 |gi|413931 |gi|42009 1 3080 1 5756 1 4896 1 4238 1 2248 1 7562 (nt) 1 730 1 860 Start (nt) 1 2024 1 1430 Contig |ORF | ID |ID | - --~ œ σ _ Ξ <u>۾</u> œ — Ξ S

Table

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	Start (nt)	Stop	match acession	match gene name	# Sim	+	length (nt)
1 132	5	4151	1 4363	gi 871048		73	43	213
140	9	5952	4324	gi 634107	kdpB [Escherichia coli]	73	59	1629
142	9	0907	1 5939	gi 410125	ribG gene product (Bacillus subtilis)	73	57	1122
149	4	1866	1717	gi 460892 	heparin binding protein-44, HBP-44 [mice, Peptide, 360 aa] pir JX0281 JX0281 heparin-binding protein-44 precursor - mouse gi1220434 ORF [Mus musculus] (SUB 2-360)	73	53	150
158		1	1431	gi 882504	ORF_f560 [Escherichia coli]	73	57	1431
1 174	9	5352	4525	9111146240	ketopantoate hydroxymethyltransferase [Bacillus subtilis]	73	55	828
175	8	5537	1 5178	gi 854657	Na/H antiporter system ORF3 [Bacillus alcalophilus]	73	56	360
1 186	5	6593	1 5493	gi 467477	unknown [Bacillus subtilis]	73	48	1101
1 249	9	6283	1 5729	gi 1524397	glycine betaine transporter OpuD (Bacillus subtilis)	73	56	555
1 265	4	1873	1 2280	gi 39848		73	41	408
1 270	7	328	1 582	gi 780461	1220 kDa polyprotein (African swine fever virus)	7.3	53	255
1 278	4	4283	3618	gi 1208965	hypothetical 23.3 kd protein [Escherichia coli]	73	49	1 999
1 279	8	4984	1 3593	gi 1185288		73	28	1392
1 291	4	1207	1 1575	gi 1511440	glutaminefructose-6-phosphate transaminase [Methanococcus jannaschii]	73	63	369
1 299	2	735	11166	gi 467437	unknown (Bacillus subtilis]	73	58	432
1 299	5	1 2050	3234	gi 467439	temperature sensitive cell division [Bacillus subtilis]	73	53	1185
1 334		1237	1 728	gi 536655	ORF YBR244w [Saccharomyces cerevisiae]	73	43	510
336	7	1827	1036	1911790943	urea amidolyase [Bacillus subtilis]	73	51	792
1 374	. 3	1389	1874	gi 1405451	YneJ [Bacillus subtilis]	73	55	486 1
433	4	1916	1 2554	gi 473902	[alpha-acetolactate synthase [Lactococcus lactis]	73	54	639
1 509	7	1795	1028	gi 467483	unknown [Bacillus subtilis]	73	56	1 89 1
1 513	-	1709	918	gi 1146220	IMAD+ dependent glycerol-3-phosphate dehydrogenase [Bacillus subtilis]	73	56	792
533	- 5	1 239	733	gi 1510605	hypothetical protein (SP:P42297) [Methanococcus jannaschii]	73	44	495
1 546	7	1148	2815	gi 41748	hsdM protein (AA 1-520) [Escherichia coli]	73	52	1668
1 549		1 762	1 382	gi 1314847	CinA [Bacillus subtilis]	73	57	381
1 567		1346	675	gi 410137	[ORFX13 [Bacillus subtilis]	73	58 1	672

Table ?

S. aureus - Putative coding regions of novel proteins similar to known proteins

	(nt)	(nt) 1112 677 209 402	acession 	exodeoxyribonuclease [Bacillus subtilis]	73	26 -	(nt) + 459
	20 20 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9		gi 1256623	[Bacillus	73	56	459
	2009						,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
	500	! ! !	gi 142010 	Shows 70.2% similarity and 48.6% identity to the EnvM protein of almonella typhimurium [Anabaena sp.]	73	57	675
1 2 1 8 2 1	209 -		gi 409286	bmrU [Bacillus subtilis]	73	52	207
3 1 2 3 1 2 3	209		gi 143320	[gap] gene products [Bacillus megaterium]	73	56	402
T E 2 T	209		gi 1063246	low homology to P14 protein of Heamophilus influenzar and 14.2 kDa protein of Escherichia coli [Bacillus subtilis]	73	56	312
m 0 H	603	911	gi 853754	ABC transporter [Bacillus subtilis]	73	28	606
	339	949	gi 143786	tryptophanyl-tRNA synthetase (EC 6.1.1.2) [Bacillus subtilis] pirlJT0481 YWBS tryptophantRNA ligase (EC 6.1.1.2) - Bacillus ubtilis	73	51	261
		3097	gi 41748	hsdM protein (AA 1-520) [Escherichia coli]	73	52	1743
		400	gi 886906	argininosuccinate synthetase [Streptomyces clavuligerus] pirIS57659 S57659 argininosuccinate synthase (EC 6.3.4.5) - treptomyces clavuligerus	73	65	399
857 11 3	e	290	gi 348052	acetoin utilization protein (Bacillus subtilis)	73	50	288
1008 + 1 79	1 062	398	gi 40100	rodc (tag3) polypeptide (AA 1-746) [Bacillus subtilis] ir S06049 S06049 rodc protein - Bacillus subtilis p P13485 TAGF_BACSU TECHOIC ACID BIOSYNTHESIS PROTEIN F.	73	41	393
1018 1 1		213	gi 529357	No definition line found [Caenorhabditis elegans] sp P46975 STT3_CAEEL OLIGOSACCHARYL TRANSFERASE STT3 SUBUNIT OMOLOG.	73	53	213
1 1033 1 3	3	491	gi 142706	comGl gene product [Bacillus subtilis]	73	51	489
1174 1 39	395	204	gi 1149513	alpha3a subunit of laminin 5 [Homo sapiens]	73	1 09	192
1175 1 65	655	329	911473817	'ORF' (Escherichia coli]	73	57	327
1187 1 3	e	209	gi 580870	ipa-37d goxA gene product (Bacillus subtilis)	73	52	207
1206 1 1 7	72	245	gi 144816	formyltetrahydrofolate synthetase (FTHFS) (ttg start codon) (EC .3.4.3) [Hoorella thermoacetica]	73	43	174
1454 1 42	423	241	gi 1213253	unknown [Schizosaccharomyces pombe]	73	53	183
1469 1 51	517	260	gi 1303787	YqeG [Bacillus subtilis]	73	55	258
	374	189		Mst26Aa gene product [Drosophila simulans]	73 1	34	186
1 1849 1 46	467	243	1911162307	DNA topoisomerase II [Trypanosoma cruzi]	73 (09	225
1 2055 1 2	2	400	gi 559381	P47K protein [Rhodococcus erythropolis]	73	34	399
2556 1 2	2	244	gi 145925	[fecB [Escherichia coli]	73	62	243

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start Start (nt)	Stop (nt)	match acession	match gene name	% sim	* ident	length (nt)
2947	1 2	549	1 400	gi 1184680	polynucleotide phosphorylase (Bacillus subtilis)	73	51	150
1 2956		746	375	gi 143397	quinol oxidase (Bacillus subtilis)	73	58	372
1 3037		655	329	gi 143091	acetolactate synthase [Bacillus subtilis]	73	55	327
3115		385	1 194	gi 323866	overlapping out-of-phase protein [Eggplant mosaic virus] sp1820129 V70K_EPWV 70 KD PROTEIN.	73	53	192
1 3603	7	1 700	1 527	gi 1439521	glutaryl-CoA dehydrogenase precursor [Mus musculus]	73	48	174
3743		798	400	gi 450688	hsdW gene of Ecopril gene product [Escherichia coli] pir[S394371838437 hsdM protein - Escherichia coli pir S09629 S09629 hypothetical protein A - Escherichia coli {S0B 40-520}	73	5.42	399
3752	1	640	1 359	gi 1524193	unknown [Mycobacterium tuberculosis]	73	59	282
3852		2	181	gi 216746	D-lactate dehydrogenase [Lactobacillus plantarum]	73	1 89	180
3914	1	475	1 239	pir S13490 S134	Hydroxymethylglutaryl-CoA synthase (EC 4.1.3.5) - Chicken (fragment)	73	53	237
3914	1 2	570	343	gi 528991	unknown (Bacillus subtilis)	73	38	228
1 4069		2	316	gi 40003	oxoglutarate dehydrogenase (NADP+) (Bacillus subtilis] p P23129 ODOL BACSU 2-OXOGLUTARATE DEHYDROGENASE E1 COMPONENT (EC 2.4.2) (ALPHA- KETOGLŪTARATE DEHYDROGENASE).	73	55	315
4165	1	715	1 365	gi 1439521	glutaryl-CoA dehydrogenase precursor [Mus musculus]	73	48	351
4196	1	п	771	gi 809660	deoxyribose-phosphate aldolase (Bacillus subtilis) pir/849455/849455 deoxyribose-phosphate aldolase (EC 4.1.2.4) - acillus subtilis	73	09	177
4202		572	1 378	gi 528991	unknown [Bacillus subtilis]	73	38	195
4314		2	193	gi 436797	N-acyl-L-amino acid amidohydrolase [Bacillus stearothermophilus] sp P37112 AMA_BACST N-ACYL-L-AMINO ACID AMIDOHYDROLASE (EC .5.1.14) (AMINOACYLASE).	73	47	192
1 4393		8	1 263	gi 216267	ORF2 [Bacillus megaterium]	73	47	261
1 35	2	903	1 1973	gi 1146196	phosphoglycerate dehydrogenase [Bacillus subtilis]	72	53	1071
38	122	119094	117877	gi 602031 	similar to trimethylamine DH [Mycoplasma capricolum] pir[849950 849950 probable trimethylamine dehydrogenase (EC .5.99.7) - Mycoplasma capricolum (SGC3) (fragment)	72	54	1218
38	123	18134	119162	gi 413968	ipa-44d gene product [Bacillus subtilis]	72	54	1029
44	119	11895	112953	gi 516272	unknown [Bacillus subtilis]	72	49	1059
48	1 7	6248	7117	gi 43499	pyruvate synthase [Halobacterium halobium]	72	49	870
50	1 7	6563	5691	gi 1205399	proton glutamate symport protein [Haemophilus influenzae]	72	53	873

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	e Ein	* ident	l length
53	6	110521	9259	gi 1303956	YqjE [Bacillus subtilis]	72	52	1263
1 56	123	129549	129995	gi 467471	unknown [Bacillus subtilis]	72	47	447
69	-	1 5298	4123	gi 1354775	pfos/R [Treponema pallidum]	72	46	1176
69	- 5	4377	1 4982	gi 904198	hypothetical protein [Bacillus subtilis]	72	43	909
73		7	1 856	gi 142997	glycerol uptake facilitator (Bacillus subtilis]	72	59	855
86	113	9371	110258	gi 467435	unknown [Bacillus subtilis]	72	50	888
127	 	: 	1593	 gi 217144 	alanine carrier protein [thermophilic bacterium PS3] pir A45111 A45111 alanine transport protein - thermophilic acterium PS-3	12	56	1593
131	 	5197	2600	gi 153952 	polymerase III polymerase subunit (dnaE) [Salmonella typhimurium) pir A45915 A45915 DNA-directed DNA polymerase (EC 2.7.7.7) III lpha chain - Salmonella typhimurium	72	53	2598
141	4 -	1040	1978	gi 1405446		72	54	626
1 149	8 -	1 2819	1 2535	gi 606234		72	44	285
149	117	1 5472	1 5245	gi 1304472	IDNA polymerase [Unidentified phycodnavirus clone OTU4]	72	55	1 228
154			1 210	gi 1205620		72	40	210
155		2207	1320	gi 391610 	[farnesy] diphosphate synthase [Bacillus stearothermophilus] pir JX0257 JX0257 geranyltranstransferase (EC 2.5.1.10) - Bacillus tearothermophilus	72	57	8888
1 180		7	328	gi 433630	A180 [Saccharomyces cerevisiae]	72	62	327
1 184	e 	1145	1 3553	gi 1205110	virulence associated protein homolog (Haemophilus influenzae)	72	49	2409
1 195	7	1 1923	1279	gi 1001730	hypothetical protein [Synechocystis sp.]	72	45	645
1 206	113	114646	115869	gi 1064807	ORTHININE AMINOTRANSFERASE [Bacillus subtilis]	72	50	1224
1 209	7	1 462	932	gi 1204666	[hypothetical protein (GB:X73124_53) [Haemophilus influenzae]	72	09	471
215	7	1 764	522	gi 881513 	insulin receptor homolog (Drosophila melanogaster) pir S57245 S57245 insulin receptor homolog - fruit fly (Drosophila elanogaster) (SUB 46- 2146)	72	63	243
1 224		2	1 790	gi 949974	sucrose repressor [Staphylococcus xylosus]	72	54	789
1 233		1526	1 765	gi 1408493		72	52	762
1 240		1 220	1 1485	gi 537049	ORF_0470 [Escherichia coli]	72	52	1266
1 245	-		1340	gi 1204578	Inypothetical protein (GB:U06949_1) [Haemophilus influenzae]	72	46	1338
	+1.	1	+				: : : : :	

237

| % sim | % ident | length (nt) 53 54 49 20 44 53 70 26 52 49 54 55 52 53 52 52 55 49 48 26 28 72 72 72 72 72 72 72 72 72 72 72 72 72 72 72 72 |44% identical amino acids with the Escherichia coli smba supress; putative IThe polymorphysm (RFLP) of this gene is associated with usceptibility to essential hypertension. The SA gene product has light homology to acetyl-NADPH-sulfite reducatase flavoprotein component (Salmonella yphimurium) | pir|S34312|S34312 hypothetical protein V - Clostridium cetobutylicum |UDP-N-acetylglucosamine 1-carboxyvinyltransferase [Bacillus ubtilis] S. aureus - Putative coding regions of novel proteins similar to known proteins glutamine-binding periplasmic protein [Haemophilus influenzae] |highly conserved among eubacteria [Clostridium acetobutylicum] |ferrous iron transport protein B [Methanococcus jannaschii] |hypothetical protein (SP:P27857) [Haemophilus influenzae] |protein-export membrane protein [Haemophilus influenzae] |virulence-associated protein E [Dichelobacter nodosus] | 2654 | 2505 |pir|S00601|BXSA |antibacterial protein 3 - Staphylococcus haemolyticus (ATP synthase subunit epsilon [Bacillus subtilis] [ABC transporter subunit [Cyanophora paradoxa] |unknown protein [Azotobacter vinelandii] (citrate synthase II (Bacillus subtilis) |orfRM1 gene product [Bacillus subtilis] (ceuD gene product [Campylobacter coli) [protein-dependent [Bacillus subtilis] [quinol oxidase [Bacillus subtilis] CoA synthetase [Homo sapiens] [putative [Bacillus subtilis] |ORF1 [Staphylococcus aureus] [unknown [Bacillus subtilis] IORF2 [Synechococcus sp.] |YrkE [Bacillus subtilis] [YqgS [Bacillus subtilis] [Bacillus subtilis] match gene name acession |gi|1340128 gi | 1205330 |gi|1070015 |gi|1204445 1gi11510643 |gi|1402944 |gi|1016162 1gi11303704 |gi|1107530 1gi11303866 |gi|1146214 |gi|1146183 |gi|1377842 1gi11088269 |gi|312379 |gi|1204497 |gi|666014 |gi|310859 |gi|153929 1911853767 |gi|143399 1gi1487433 |gi|433992 gi|563258 9308 1245 1609 110493 1263 3466 1184 1094 5039 625 1251 1 746 573 1 957 1 435 432 1 260 592 1 788 421 2177 3047 (nt) | 1 | 1142 1 2108 1 5326 1 9622 1 7 |12538 1 1 3253 1086 1 285 | 517 926 340 81 295 295 1 1 840 1 1 1 150 1 1 1 130 | 1 | 2 ლ — 7 11 | 1 Contig |ORF | ID |ID | 7 - - 2 110 11 <u>-</u> 7 9 ~ 7 460 259 304 315 316 337 364 409 453 463 519 559 671 619 687 837 868 307

810

288 258 315 684 2046

924 588 150 624 1626 1104

420 501 744

570

591

957

435 639 303

Table

S. aureus - Putative coding regions of novel proteins similar to known proteins

							*	# ! ! ! ! !
Contig ID	g lorf	Start (nt)	Stop (nt)	match acession	match gene name	mis &	% ident	length (nt)
1209	-	383	213	gi 144735	neurotoxin type B [Clostridium botulinum]	72	44	171
1469	7	671	474		hypothetical protein (GB:D26562 47) [Haemophilus influenzae]	127	63	198
1956		127	365	gi 154409 	hexosephosphate transport protein (Salmonella typhimurium) pir[B4]853 B41853 hexose phosphate transport system regulatory rotein uhpB - Salmonella typhimurium	72	4	363
1 2101		e 	401	gi 1303950	YqiY [Bacillus subtilis]	72	50	399
1 2503	·	1 569	399	gi 149713 	formate dehydrogenase [Methanobacterium formicicum] pir A42712 A42712 formate dehydrogenase (EC 1.2.1.2) - ethanobacterium formicicum	72	56	171
1 2967		e 	155	gi 1212729	YqhJ [Bacillus subtilis]	127	46	153
3004		1 367	1 185		hypothetical protein [Bacillus subtilis]	72	55	183
1 3109		1 278	141	gi 413968		72	45	138
3171		m 	1 287	gi 515938 	glutamate synthase (ferredoxin) [Synechocystis sp.] pir S46957 S46957 glutamate synthase (ferredoxin) (EC 1.4.7.1) - ynechocystis sp.	72	52	285
3771		1 26	1 367	gi 1408501 	homologous to N-acyl-L-amino acid amidohydrolase of Bacillus stearothermophilus [Bacillus subtilis]	72	63	342
3951			1 222	gi 1500409	[M. jannaschii predicted coding region MJ1519 [Methanococcus jannaschii]	72	38	222
4190		121	1 362	gi 39956		72	57	360
1 4444		e 	347	gi 1009366	Respiratory nitrate reductase [Bacillus subtilis]	72	55	345
9	- 5	931	1200	lgi1537095	ornithine carbamoyltransferase [Escherichia coli]	71	55	270
=	115	111350	110859	gi 532309	155 kDa protein [Escherichia coli]	71	47	492
19	- 5	1248	1 2435	gi 1244574	D-alanine:D-alanine ligase [Enterococcus hirae]	71	52	1188
1 21		868	1488	gi 149629 	anthranilate synthase component 2 [Leptospira biflexa] pirlC328401C32840 anthranilate synthase (EC 4.1.3.27) component II Leptospira biflexa	71	45	591
34	-		1 567	gi 1303983	YqkF [Bacillus subtilis]	71	59	567
1 37	e 	3192	1 2806	gi 1209681	glutamate-rich protein [Bacillus firmus]	71	50	387
38	- 18	112250	112462	gi 927645	arginyl endopeptidase (Porphyromonas gingivalis)	71	50	213
39		1246	4431	pir S09411 S094	r S09411 S094 spoiliE protein - Bacillus subtilis	71	49	3186
53	114	115770	114760	gi 142611	branched chain alpha-keto acid dehydrogenase El-alpha [Bacillus ubtilis]	71	58	1011
54	111	13461	112625	gi 143014	gnt repressor [Bacillus subtilis]	71	46	837

Table 2

1125 306 699 1110 531 384 315 966 1365 441 549 336 810 849 174 699 576 525 438 462 963 879 | % ident | length 47 41 47 53 49 45 57 40 53 51 52 26 49 46 20 54 48 20 48 99 54 38 7.1 71 71 8 sim |high homology to flavohemoprotein (Haemoglobin-like protein) of Alcaligenes |EIIC domain of PTS-dependent Gat transport and phosphorylation Escherichia | eutrophus and Saccharomyces cerevisiae (Bacillus subtilis) [probable protein kinase [Saccharomyces cerevisiae] |repressor protein [Streptococcus sobrinus] |membrane bound protein [Bacillus subtilis] |hypothetical protein [Bacillus subtilis] lipa-61d gene product [Bacillus subtilis] |hypothetical protein (Bacillus subtilis] |hypothetical protein (Bacillus subtilis) 120 kDa protein (Streptococcus gordonii) Opine dehydrogenase [Arthrobacter sp.] |ORF4 gene product [Bacillus subtilis] |unknown [Mycobacterium tuberculosis] liep protein [Bacillus subtilis] |phnE protein [Escherichia coli] |putative [Bacillus subtilis] |GlnQ [Mycoplasma pneumoniae] ORF1 [Campylobacter jejuni] [HisIE [Lactococcus lactis] |G40P [Bacteriophage SPP1] ORF210 [Escherichia coli] [YqiG [Bacillus subtilis] [YqhX [Bacillus subtilis] [YqeH [Bacillus subtilis] | match gene name | 13897 | 114334 | gi| 1063247 |gi|1303926 (gi11403403 | 6 | 3322 | 2771 |gi|1256625 |gi|1060848 |gi|1303788 |gi|1303913 |gi|1215694 |gi|147198 |gi|508175 |gi|143089 |gi|413985 1gi1663254 |gi|143156 |gi|425488 |gi|633732 | 2273 |gi|709993 | 1496 | gi|143035 |gi|310634 |gi|904205 |gi|529650 |gi|149384 | 5 | 3349 | 2720 |gi|303562 19i1709991 |gi|49318 3087 17152 | 5860 11234 1 9831 (10955 1 2394 | 2089 115695 1723 111689 1 2745 1 5169 Contig |ORF | Start | Stop ID | ID | (nt) | (nt) 5895 2022 8269 316 532 1 8505 115312 6598 3635 111 111354 1936 4 | 1849 2848 17 1 7601 1 1 1 1062 1 3087 1743 17 1 4501 1 4822 6335 1 1 630 1 272 1 2 | 618 1 1 1 136 116 7 118 112 -9 118 ~ 119 -5 ~ -114 113 164 169 57 62 98 96 100 103 109 133 154 193 233

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	* sim	% ident	length (nt)
301	9	3492	2461	gi 467417	similar to lysine decarboxylase [Bacillus subtilis]	71	57	1032
306	4	6607	5222	gi 1256618	transport protein [Bacillus subtilis]	71	56	1386
307	2	1536	925	gi 602683	orfC [Mycoplasma capricolum]	71	45	612
310	5	5793	5146	gi 348052	Jacetoin utilization protein [Bacillus subtilis]	71	51 1	648
1 322		7	1303	gi 1001819	hypothetical protein [Synechocystis sp.]	71	46	1302
1 333	4	4171	3995	gi 467473	unknown [Bacillus subtilis]	71	57	1771
350	7	548	922	gi 551879	ORF 1 [Lactococcus lactis]	71	55	375
375	4	1860	3071	gi 467447	unknown [Bacillus subtilis]	71	57 1	1212
380	2	1560	2102	lgi 142557	ATP synthase b subunit [Bacillus megaterium]	71	43	543
414	2	251	637	gi 580904	homologous to E.coli rnpA (Bacillus subtilis)	71	49	387
424		335	1354	gi 581305	L-lactate dehydrogenase [Lactobacillus plantarum]	71 1	57	1020
436	4	3701	3270	pir PN0501 PN05	phosphoribosylanthranilate isomerase (EC 5.3.1.24) - Bacillus subtilis (fragment)	71	99	432
1 482	1	ъ	1280	gi 410142	ORFX18 [Bacillus subtilis]	71	49	1278
525	e 	2272	1844	gi 143370 	phosphoribosylpyrophosphate amidotransferase (PUR-F; EC 2.4.2.14) Bacillus subtilis}	71	56	429
529	4	2739	1 2047	gi 606150	ORF_f309 [Escherichia coli]	71	43	1 869
563	7	22	696	gi 1237015	ORF4 [Bacillus subtilis]	71	53	948
581		506	255	gi 1301730	[12563.2 [Caenorhabditis elegans]	71	47	252
612	2	1068	913	gi 153968	fimbriae Z [Salmonella typhimurium]	71	55	156
613	. –	-	654	, gi 466778	lysine specific permease [Escherichia coll]	71	50	654
618	FH -	1243	623	gi 1146238	poly(A) polymerase [Bacillus subtilis]	71	52	621
630		1170	586	gi 1486243	unknown [Bacillus subtilis]	71	53	585
691		1126	641	gi 289260	comE ORF1 [Bacillus subtilis]	71	51	486
694	2	149	427	gi 12971 	NADH dehydrogenase subunit V (AA 1-605) (Gallus gallus) ir[S10197]S10197 NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain - chicken mitochondrion (SGC1)	71	47	279
1 715	1 2	169	777	gi 1303830	YqfL [Bacillus subtilis]	71	53	1 609
746	- 5	1473	076	gi 1377843	unknown [Bacillus subtilis]	71	52	504

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

+	+-	+			· · · · · · · · · · · · · · · · · · ·	+	+	+
Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
1 748		1437	1 802	gi 1405459	YneS (Bacillus subtilis)	71	649	636
1 753		1018	1 524	gi 1510389	M. jannaschii predicted coding region MJ0296 [Methanococcus jannaschii]	71	53	495
1 761	-	ю	1 215	gi 475972	pentafunctional enzyme [Pneumocystis carinii]	71 +	47	213
1 783		1203	1 703	gi 536655	ORF YBR244w [Saccharomyces cerevisiae]	71	52	501
008 1		1292	1 987	gi 1204326		71	48	306
908	-	116	1 286	gi 1419075	cbiM gene product [Methanobacterium thermoautotrophicum]	71	20	171
931		973	488	gi 893358	PgsA (Bacillus subtilis)	71	26	486
1041		7	1 262	gi 1408507	pyrimidine nucleoside transport protein (Bacillus subtilis)	71	45	261
1070		2	172	lgi1709993	hypothetical protein [Bacillus subtilis]	7.1	46	171
1176		57	365	gi 151259	HMG-COA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir A44756 A44756 hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp.	11	49	309
1181	-	366	184	gi 46971	epiP gene product [Staphylococcus epidermidis]	71	20	183
1 1281		e -	1 290	gi 153016	ORF 419 protein [Staphylococcus aureus]	71	20	288
1348	-	456	1 229	gi 602683	orfC [Mycoplasma capricolum]	71	48	228
2002		756	1 379	gi 1008177	ORF YJL046w [Saccharomyces cerevisiae]	71	48	378
2119		7	1 217	gi 1046088	arginyl-tRNA synthetase [Mycoplasma genitalium]	71	50	216
2418		£	1 320	gi 1499771	M. jannaschii predicted coding region MJ0936 [Methanococcus jannaschii]	71	57	318
1 2961		7	1 187	gi 312443	carbamoyl-phosphate synthase (glutamine-hydrolysing) [Bacillus aldolyticus]	71	57	186
2999	- 5	1 67	306	gi 710020	nitrite reductase (nirB) [Bacillus subtilis]	71	43	240
3033		2	1 184	gi 1262335	YmaA [Bacillus subtilis]	71	57	183
3584		т -	1 338	gi 401716	beta-isopropylmalate dehydrogenase [Neurospora crassa]	71	55	336
3715	2	743	399	gi 563952		71	1 65	345
3785	H	1 770	1 387	gi 47382	acyl-CoA-dehydrogenase [Streptomyces purpurascens]	71	57	384
3875		541	1 272	gi 1001541	Inypothetical protein [Synechocystis sp.]	71	38	270
4135		637	320	gi 142695 	S-adenosyl-L-methionine:uroporphyrinogen III methyltransferase Bacillus megaterium]	71	52	318
4249		. 63	239	gi 1205363	deoxyribose aldolase [Haemophilus influenzae]	71	63	177
1 4508		1 530	1 267	gi 1197667	vitellogenin [Anolis pulchellus]	71	46	264
1	 					-		-

Table 7

S. aureus - Putative coding regions of novel proteins similar to known proteins

	ORF	+	Stop	-+	match gene name	# 8 sim	+	
	OI.	(nt)	(nt)	acession			-	(nt)
9	<u>ش</u>	1237	1 2721	gi 1321788	arginine ornithine antiporter [Clostridium perfringens]	10	54	1485
11	111	6572	7486	gi 216854	P47K [Pseudomonas chlororaphis]	10	41	915
12		1 2890	1481	gi 467330	replicative DNA helicase (Bacillus subtilis]	07	49	1410
1 15		1756	1 893	gi 451216	Mannosephosphate Isomerase [Streptococcus mutans]	70	46	864
1 15	- 5	7721	1 1050	gi 476092	unknown [Bacillus subtilis]	70	20	228
1 17	7	2132	1350	gi 145402	choline dehydrogenase [Escherichia coli]	0,4	52	783
21		7	925	gi 149516	anthranilate synthase alpha subunit [Lactococcus lactis] pir[S35124 S35124 anthranilate synthase (EC 4.1.3.27) alpha chain - actococcus lactis subsp. lactis	70	20	924
1 25	7	1 5580	6251	gi 1389549	ORF3 (Bacillus subtilis)	07	1 52	672
1 33	9	6071	7423	gi 1303875	YqnB (Bacillus subtilis)	07	1 51	1353
36	7	959	1594	gi 500755	methyl purine glycosylase [Mus musculus]	10	47	989
38	8	4901	1 5860	gi 1408507	pyrimidine nucleoside transport protein [Bacillus subtilis]	70	1 44	096
1 44	80	1 5312	2989	gi 1006620	hypothetical protein (Synechocystis sp.]	07	49	678
46	10	0568	110020	gi 1403126	czcD gene product [Alcaligenes eutrophus]	10	1 45	1071
52	1 5	7272	1900	gi 1486247	unknown [Bacillus subtilis]	10	53	828
52	· •	4048	1 4656	gi 244501 	esterase II=carboxylesterase (EC 3.1.1.1) [Pseudomonas fluorescens, eptide, 218 aa]	70	20	609
1 56	8	1 8460	1 9962	gi 1339951	small subunit of NADH-dependent glutamate synthase [Plectonema boryanum]	10	1 51	1503
1 62		48	1 290	gi 142702	A competence protein 2 (Bacillus subtilis)	10	1 47	243
1 64		1080	1 541	gi 1204377	molybdopterin biosynthesis protein [Haemophilus influenzae]	1 70	1 47	540
100	2	5139	3595	gi 1204834	[2',3'-cyclic-nucleotide 2'-phosphodiesterase [Haemophilus influenzae]	10	1 47	1545
91	4	1 7793	5466	gi 886471	methionine synthase [Catharanthus roseus]	1 70	56	2328
96	2	8754	7255	pir B39096 B390	alkaline phosphatase (EC 3.1.3.1) III precursor - Bacillus subtilis	07	1 54	1500
1 110	7	191	1300	gi 145294	adenine phosphoribosyl-transferase [Escherichia coli]	10	51	534
116	9	7026	1 7976	gi 143607	sporulation protein (Bacillus subtilis)	07	1 50	951
121	8	6401	1 6988	gi 1107528	ttg start [Campylobacter coli]	10	45	588
131	B 	6842	1 7936	gi 1150454	prolidase PepQ [Lactobacillus delbrueckii]	10	1 48	1095
	·	1 +	· · · · · · · · · · · · · · · · · · ·	111111111111111111111111111111111111111				

| length (nt) | % ident | & sim 7.0 putative membrane-bound protein with four times repitition of ro-Ser-Ala at open reading frame; putative [Bacillus amyloliquefaciens] pir/B29091|B29091 hypothetical protein (bglA region) - Bacillus myloliquefaciens (fragment) peptide transport system protein SapF homolog; SapF homolog [Mycoplasma phosphoribosyl glycinamide formyltransferase (PUR-N) [Bacillus ubtilis] S. aureus - Putative coding regions of novel proteins similar to known proteins |pir|C38530|C385 |queuine tRNA-ribosyltransferase (EC 2.4.2.29) - Escherichia coli |high affinity ribose transport protein [Haemophilus influenzae] |hypothetical protein (GB:X73124_26) [Haemophilus influenzae] |hypothetical protein (GB:D10483_18) [Haemophilus influenzae] hypothetical protein (SP:P37002) [Methanococcus jannaschii] the N-terminus; function unknown [Alcaligenes utrophus] |ATP synthase gamma subunit [Bacillus megaterium] |ruvA protein (gtg start) [Escherichia coli] aminopeptidase a/i [Haemophilus influenzae] (hypothetical protein (Bacillus subtilis) |uracil permease [Bacillus caldolyticus] |ORF1 gene product [Bacillus subtilis] |protein-dependent (Bacillus subtilis) [excisionase [Bacteriophage 154a] [aspartokinase II [Bacillus sp.] |GumF [Xanthomonas campestris] |RCH2 protein (Brassica napus) IORF1 [Staphylococcus aureus] |unknown [Bacillus subtilis] ORFF (Bacillus subtilis) |YqgZ [Bacillus subtilis] [MgtE [Bacillus firmus] | match gene name acession |gi|1205212 |gi|1215695 |gi|1340128 |gi|1070013 |gi|1204752 |gi|1511524 lgi11303873 match |gi|1204665 |gi|1205934 |gi|147782 1gi1311309 1gi 1904181 |gi|215098 |gi|142540 1gi | 733147 |gi|619724 |gi|727145 |gi|602292 |gi|881434 1gi | 467340 |gi|142560 |gi|143372 |gi|431231 1gi149315 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 1 3827 699 | 1 417 1 703 1 3201 1 1683 1 1 1 1383 1 1 830 1 727 ~ ~ <u>ო</u> <u>س</u> œ — <u>س</u> -~ ~ --

 Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

	1 4 6		4	-+	match done name	+	9	+
QI I		(nt)		acession				(nt)
681		1488	781	gi 1001678	hypothetical protein [Synechocystis sp.]	70	53	1 80 /
108		- 5	448	sp P33940 YOJH_	HYPOTHETICAL 54.3 KD PROTEIN IN ECO-ALKB INTERGENIC REGION.	107	51	447
725		51	722	gi 1001644	hypothetical protein [Synechocystis sp.]	70	4.8	672
1 776		1371	187	gi 145165	putative [Escherichia coli]	1 07	47	585
834		1 250	1 783	gi 552971	NADH dehydrogenase (ndhF) [Vicia faba]	107	47	534
865	- 5	1585	1379	gi 1204636	ATP-dependent helicase [Haemophilus influenzae]	1 07	45	207
1 894		535	1 269	gil467364	(DNA binding protein (probale) [Bacillus subtilis]	107	41	267
919		m	317	gi 1314847	CinA [Bacillus subtilis]	70	40	315
1 944		<u>ش</u>	572	1911709991	hypothetical protein [Bacillus subtilis]	70	44	570
886	7	1772	605	gi 142441	ORF 3; putative [Bacillus subtilis]	1 07	20	168
1055	п	ю —	335	gi 529755	spec [Streptococcus pyogenes]	107	37	333
1 1093		7	904	gi 853754	ABC transporter [Bacillus subtilis]	70	49	903
1 1109		7	310	gi 1001827	Inspothetical protein [Synechocystis sp.]	0,	42	309
1220	-	1 468	1 235	pir S23416 S234	lepiB protein - Staphylococcus epidermidis	70	40	234
1279		73	348	gi 153015	FemA protein [Staphylococcus aureus]	70	47	276
1336		195	542	sp P31776 PBPA_	PENICILLIN-BINDING PROTEIN IA (PBP-1A) (PENICILLIN-BINDING PROTEIN A).	107	50	348
1537	1 2	232	1 402	gi 1146181		107	50	171
1574	1	451	272	gi 219630	endothelin-A receptor (Homo sapiens)	70	47	180
1640		069	346	gi 1146243 	122.4% identity with Escherichia coli DNA-damage inducible protein; putative (Bacillus subtilis)	70	46	345
2504		2	1 286	gi 495179	transmembrane protein [Lactococcus lactis]	70	51	285
3061		564	1 301	gi 508175 	EIIC domain of PTS-dependent Gat transport and phosphorylation Escherichia coli	70	44	264
3128	-	- 2	199	gi 1340096	unknown [Mycobacterium tuberculosis]	70	51	198
3218		т 	488	gi 515938 	glutamate synthase (ferredoxin) [Synechocystis sp.] pir S46957 S46957 glutamate synthase (ferredoxin) (EC 1.4.7.1) - ynechocystis sp.	7.0	50	486
3323		1 794	1 399	gi 1154891	ATP binding protein [Phormidium laminosum]	70	52	396
3679		599	1 399	gi 529385		70	30	201
-	í							

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	mis %	8 ident	l length
3841		902	398	gi 1208965	hypothetical 23.3 kd protein (Escherichia coli)	70	1 47	309
3929	-	8	401	gi 149435	putative [Lactococcus lactis]	7.0	49	399
4044		595	374	gi 602031	similar to trimethylamine DH [Mycoplasma capricolum] pir[849950 849950 probable trimethylamine dehydrogenase (EC .5.99.7) - Mycoplasma capricolum (SGC3) (fragment)	70	40	222
4329		558	1 280	gi 1339951	small subunit of NADH-dependent glutamate synthase [Plectonema boryanum]	0/	49	279
1 4422		576	289	lgi1296464	ATPase [Lactococcus lactis]	70	1 57	1 288 1
1 4647	-	361	1 200	gi 166412	NADH-glutamate synthase [Medicago sativa]	70	65	162
1 16		7571	9031	gi 1499620	M. jannaschii predicted coding region MJ0798 [Methanococcus jannaschii]	69	44	1461
1 16		0806	110033	gi 1353197	thioredoxin reductase [Eubacterium acidaminophilum]	69	1 54	954
30		1452	727	4910	hypothetical protein (GB:U14003_302) [Haemophilus influenzae]	69	1 52	726
38	4	1023	1298	gi 407773	devA gene product [Anabaena sp.]	69	1 41	276
44	5	5987	6595	gi 1205920	molybdate uptake system hydrophilic membrane-bound protein [Haemophilus influenzae]	69	45	1 609 1
62	115	9104	9475	gi 385178	unknown [Bacillus subtilis]	69	44	372
99	4	2402	2803	gi 1303893	YqhL [Bacillus subtilis]	69	1 51	402
1 69 1	115 1	114124	113627	gi 149647	ORFZ [Listeria monocytogenes]	69	1 37	498
1 67 1		114053	114382	lgi1305002	ORF f356 [Escherichia coli]	69	49	330
1 69 1	119 1	15130	115807	gi 1109684	ProV [Bacillus subtilis]	69	1 45	678
1 78	9	1447	2124	gi 1256633	putative (Bacillus subtilis)	69	53	678
1 78	4	4513	3725	gi 1303958	[YqjG [Bacillus subtilis]	69	32	1 682
85	4	4521	4213	pir E29326 E293	hypothetical protein (pur operon) - Bacillus subtilis	69	32	309
86	9	3253	2654	gi 973332	OrfC [Bacillus subtilis]	69	1 50	1 009 1
95		96	710	gi 786468	4All antigen, sperm tail membrane antigen=putative sucrose-specific i hosphotransferase enzyme II homolog [mice, testis, Peptide Partial, 72 aa]	69	43	615
1 100	7	6023	7426	gi 1205355	Na+/H+ antiporter [Haemophilus influenzae]	69	39	1404
102	2	2678	1650	gi 561690	sialoglycoprotease (Pasteurella haemolytica)	69	47	1029
1 103	8 1	112241	8537	gi 1009366	Respiratory nitrate reductase (Bacillus subtilis)	69	54	3705
103	11	114987	112552	gil710020	nitrite reductase (nirB) [Bacillus subtilis]	69	1 51	2436

| % ident | length 69 69 69 69 69 69 69 69 69 69 69 69 69 8 sim |pir|S28089|S280 |hypothetical protein A - yeast (Zygosaccharomyces bisporus) plasmid pSB3 phenylalany1-tRNA synthetase beta subunit (AA 1-804) [Bacillus btilis] | SP|P17731|HIS8 | HISTIDINOL-PHOSPHATE AMINOTRANSFERASE (EC 2.6.1.9) (IMIDAZOLE ACETOLlhexosephosphate transport protein [Salmonella typhimurium]
l pir|D41853|D41853 hexose phosphate transport system protein uhpT | almonella typhimurium |pyruvate formate-lyase activating enzyme [Haemophilus influenzae] |hypothetical protein (GB:U14003_302) [Haemophilus influenzae] |pir|A42771|A427 |reticulocyte-binding protein 1 - Plasmodium vivax |hsdR protein (AA 1-1033) [Escherichia coli] |elongation factor Ts [Spirulina platensis] |hypothetical protein [Synechocystis sp.] lipa-7d gene product [Bacillus subtilis] |FimE protein [Escherichia coli] |YneP [Bacillus subtilis] [NrdF [Bacillus subtilis] |YbbF [Bacillus subtilis] | PHOSPHATE TRANSAMINASE) lo307 [Escherichia coli] | match gene name acession |gi|1154633 |gi|1205538 |gi|1256135 |gi|1405456 |gi|1001768 | 111 | 8708 | 110168 | 9i|154411 |gi|290509 |gi|413931 |gi|732682 |gi|40054 1gi141750 8339 1 1671 5796 1537 | 1058 Contig |ORF | Start | Stop ID | ID | (nt) | (nt) 110400 1 2777 813 319 1 953 854 1598 9898 6038

(nt)

1002 2412 921 480 267 186 552 1188 2775 243 318 3492 291 231 426 693 957 1029 1041 588 348 957 48 54 20 43 53 43 47 45 49 48 20 61 45 28 40 44 39 39 52 69 48 53 30 69 69 69 69 69 69 purine synthesis repressor [Haemophilus influenzae] phosphoglyceromutase [Zymomonas mobilis] pyruvate carboxylase [Rhizobium etli] ORF [Balaenoptera acutorostrata] T04H1.4 [Caenorhabditis elegans] |arginase (Bacillus caldovelox) [mutY homolog [Homo sapiens] |unknown [Bacillus subtilis] ORF1 (Bacillus subtilis) | 16 | 16644 | 17414 | gi | 1204435 |gi|1276985 | 2 | 3286 | 2246 |gi|1458228 |gi|1221782 |gi|1122758 |gi|1256798 |gi|1491664 |gi|336458 |gi|556015 | 3 | 1210 | 1761 |gi|296031 18 | 4820 | 5776 1 6 | 4309 | 5310 1 4 | 2222 | 4633 | 7384 | 3893 1808 1 2424 7741 1129 1 3 1 2075 6713 127 |10666 1 1 | 2085 1 1 | 5551 1 3 | 2098 1999 1 2 | 33 1 1 636 828 1 2 | 267 12 | 484 1 87 - 2 -112 7 3 **в** 4 1 206 121 125 149 190 421 113 114 161 165 169 351

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	10RF	Start (nt)	Stop (nt)	match acession	match gene name	# Si ju	% ident	length (nt)
509	e -	1730	1371	gi 49224	URF 4 [Synechococcus sp.]	69	39	360 1
1 520	20	3023	2823	gi 726427	<pre>(similar to D. melanogaster MST101-2 protein (PIR:S34154) Caenorhabditis (elegans)</pre>	69	39	201
1 531		26	1 760	gi 509672	repressor protein [Bacteriophage Tuc2009]	1 69 1	33	735
689		107	1 253	gi 169101	17.9 kDa heat shock protein (hspl7.9) [Pisum sativum]	69	52	147
1 594	2	597	1391	gi 142783	IDNA photolyase [Bacillus firmus]	69	48	1 367
1 604	4	2476	2114	lgi1413930	lipa-6d gene product [Bacillus subtilis]	69	45	363
1 607		7	313	gi 1236103	W08D2.3 [Caenorhabditis elegans]	69	47	312
1 607	7	1 590	312	lgi1536715	ORF YBR275c [Saccharomyces cerevisiae]	69	39	279
734		864	1 433	gi 467327	unknown [Bacillus subtilis]	69	44	432
1 759	-	е	1 338	gi 1009367	Respiratory nitrate reductase [Bacillus subtilis]	1 69 1	50	336
761	2	392	286	gi 3508 	Leucyl-tRNA synthetase (cytoplasmic) [Saccharomyces cerevisiae] 1370340 ORF YPL160w [Saccharomyces cerevisiae]	69	46	195
1 802		72	1013	gi 143044	ferrochelatase [Bacillus subtilis]	69	55	942
816	1	1 2573	1368	gi 1510268	restriction modification system S subunit [Methanococcus jannaschii]	1 69	45	1206
838	- 5	133	387	gi 1255371	lcoded for by C. elegans cDNA yk34a9.5; coded for by C. elegans cDNA yk34a9.3; Similar to guanylate kinase [Caenorhabditis elegans]	69	46	255
851	1 2	745	1005	qi 288998	secA gene product [Antithamnion sp.]	69	39	261
1 867	-	1 535	1 269	gi 1070014	protein-dependent [Bacillus subtilis]	69	47	267
995		954	478	g1 1205569	[transcription elongation factor [Haemophilus influenzae]	69	53	477
666		1009	905	gi 899254	predicted trithorax protein [Drosophila virilis]	69	21	504
1 1127	7	1315	629	gi 1205434	H. influenzae predicted coding region HI1191 [Haemophilus influenzae]	69	56	657
1138	-	248	1 460	gi 1510646	M. jannaschii predicted coding region MJ0568 [Methanococcus jannaschii]	69	48	213
1 2928	-	ю	401	gi 290503	glutamate permease [Escherichia coli]	69	41	399
1 3090		444	1 223	gi 1204987	DNA polymerase III, alpha chain [Haemophilus influenzae]	69	36	222
3817		2	400	gi 1483199	peptide-synthetase [Amycolatopsis mediterranei]	69	45	399
1 3833		199	335	gi 1524193	unknown [Mycobacterium tuberculosis]	1 69 1	46	333
-					_	•	-	-

Table 2

1011 1110 186 333 285 519 2445 366 1530 240 597 1068 1038 1215 171 348 228 291 987 810 957 774 | % ident | length (nt) 46 44 64 59 36 43 44 49 45 48 55 47 39 46 40 43 57 55 48 46 20 45 | % sim 69 69 69 69 69 69 89 99 89 89 89 89 89 89 89 89 89 89 89 89 89 89 167 kDa Myosin-crossreactive streptococcal antigen (Streptococcus yogenes) |sp|P38038|CYSJ_ |SULFITE REDUCTASE (NADPH) FLAVOPROTEIN ALPHA-COMPONENT (EC 1.8.1.2) (SIRprecursor (aa -20 to 381) [Acinetobacter calcoaceticus] ir|A29277|A29277 IORF_o622; reading frame open far upstream of start; possible rameshift, IM. genitalium predicted coding region MG246 [Mycoplasma genitalium] lorfY 3' of comK [Bacillus subtilis, E26, Peptide Partial, 140 aa] | pir|543612|543612 hypothetical protein Y - Bacillus subtilis | sp|P40398|YHXD_BACSU HYPOTHETICAL PROTEIN IN COMK 3'REGION (ORFY) |similar to phosphotransferase system enzyme II [Escherichia coli] | sp|P32672|PTWC_ECOLI PTS_SYSTEM, FRUCTOSE-LIKE-2 IIC COMPONENT | PHOSPHOTRANSFERASE_ENZYME_II, C_COMPONENT). |restriction-modification enzyme subunit M1 [Mycoplasma pulmonis] | aldose 1-epimerase (EC 5.1.3.3) - Acinetobacter lcoaceticus | pir|S49395|S49395 HsdMl protein - Mycoplasma pulmonis |241k polyprotein [Apple stem grooving virus] | linking to previous ORF [Escherichia coli] |argininosuccinate lyase [Escherichia coli] hypothetical protein [Synechocystis sp.] [aldehyde dehydrogenase [Vibrio cholerae] lipa-43d gene product [Bacillus subtilis] |sporulation protein [Bacillus subtilis] |unknown [Saccharomyces cerevisiae] 135 kDa protein [Escherichia coli] [MalC [Streptococcus pneumoniae] |phnE protein [Escherichia coli] |ATPase [Enterococcus hirae] |ProW [Bacillus subtilis] |YqjA [Bacillus subtilis] |YqeR (Bacillus subtilis] пате match gene FRAGMENT) acession | 1193 |gi|1109685 1gi11303805 |gi|1208451 |qi|1045937 |gi|1303952 match |gi|546918 |gi|496158 |gi|396296 |gi|807973 |gi|155276 |gi|285608 |gi|147198 |gi|517205 1gi1606342 |gi|143608 | 1011 | | 1396307 |gi|413967 |gi|290642 |gi|145173 |gi|153724 |gi|38722 1 2074 3404 Stop 8415 4132 111671 | 110685 8155 (nt) 1 8772 1 2966 1150 3622 1 8638 3698 1594 400 374 333 1 230 750 293 | Start | (nt) 9889 3643 3 | 2484 1 1 2302 1 2592 6328 1 4 | 3536 7346 1899 7865 11115 1 2187 1 2666 1424 747 1 215 90 1 457 - 1 - 1 Contig |ORF ID | ID - 2 _ - -110 9 7 _ 4 æ — 7 6 m 4 -'n 7 4115 4079 4258 4317 4139 4465 12 31 40 46 48 48 53 70 83 116 129 m

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

11111	1		1					
Contig	IORE	Start (nt)	Stop (nt)	match acession	match gene name	& sim	% ident	length (nt)
132	m 	1 1867	1 2739	gi 216267	(RF2 [Bacillus megaterium]	1 68	48	873
134	7	848	1012	gi 147545		89	50	165
1 141	7	372	614	gi 872116		89	36	243
149	7	1 2454	1 2260	gi 145774		89	48	195
155	7	1776	1534	gi 216583	ORF1 [Escherichia coli]	89	36	243
158		1 1826	3289	sp P33940 YOJH_	HYPOTHETICAL 54.3 KD PROTEIN IN ECO-ALKB INTERGENIC REGION.	1 89 1	51	1464
169	9	1 2749	3318	gi 1403402	unknown [Mycobacterium tuberculosis]	99	46	570
175	110	9158	7365	gi 1072395	IphaA gene product (Rhizobium meliloti)	89	51	1794
188	۲ .	4184	5434	gi 1173843	13-ketoacyl-ACP synthase II [Vibrio harveyi]	89	48	1251
1 189	e 	1 907	1665	gi 467383	IDNA binding protein (probable) (Bacillus subtilis)	89	55	1 259
1 206	2	1 7683	6029	gi 1256138	YbbI (Bacillus subtilis)	89	48	975
1 206	8 -	110425	112176	lgi 452687		68	48	1752
1 212	ω. 	3421	3648	gi 1369941	cl gene product [Bacteriophage Bl]	89	39	228
1 214	8	1 5457	1 6482	lgi11420467	ORF YOR196c [Saccharomyces cerevisiae]	1 68	45	1026
1 237	4	1 2507	3088	gi 149381	HisH [Lactococcus lactis]	1 89	46	582
1 243	2	1 5540	1 4542	gi 1235684	mevalonate pyrophosphate decarboxylase [Saccharomyces cerevisiae]	1 89	47	1 666
1 262	1	m —	164	gi 150974	4-oxalocrotonate tautomerase [Pseudomonas putida]	1 89	42	162
1 262	- 1	1984	1118	gi 1147744	PSR [Enterococcus hirae]	89	49	867
1 276	9 -	3702	3139	sp P30750 ABC_E	p P30750 ABC_E ATP-BINDING PROTEIN ABC (FRAGMENT).	68	50	564
306	9	6345	5725	gi 1256617	adenine phosphoribosyltransferase [Bacillus subtilis]	1 89	53	621
333	. 3	4599	1 3850	lgi1467473	unknown [Bacillus subtilis]	1 89	45	750
1 365	9 -	5017	4838	gi 1130643	[12283.3 [Caenorhabditis elegans]	89	45	180
376	- 5	549	1646	gi 1277026		1 89	51	1098
1 405		1 1741	1 872	lgi11303917	YqiB (Bacillus subtilis)	1 89 1	47	870
1 406	- 5	1 853	539	gi 1511513	ABC transporter, probable ATP-binding subunit [Methanococcus jannaschii]	69	44	315
1 426	9	3558	1 3391	gi 624632	GltL [Escherichia coli]	1 69	48	168
1 438		108	329	gi 146923	Initrogenase reductase [Escherichia coli]	89	43	222
1			·	1	++++++++++++++++++++++++++++++++++++++	+		+

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	aris ar	% ident	length (nt)
1 443		1 476	1 240	gi 535810	hippuricase {Campylobacter jejuni]	89	42	237
1 443	7	518	1015	gi 1204742	H. influenzae predicted coding region HI0491 [Haemophilus influenzae]	89	48	498
1 443		1 4447	3779	gi 809660	deoxyribose-phosphate aldolase (Bacillus subtilis) pir S49455 S49455 deoxyribose-phosphate aldolase (EC 4.1.2.4) - acillus subtilis	89	55	699
476	~	240	1184	gi 971345	unknown, similar to E.coli cardiolipin synthase [Bacillus subtilis] sp P45860 YWIE_BACSU HYPOTHETICAL 58.2 PROTEIN IN NARI-ACDA NTERGENIC REGION.	89	24	945
486	- 5	1876	1 1046	gi 147328	transport protein [Escherichia coli]	89	41	831
517	m 	1764	1 2084	gi 1523809	orf2 [Bacteriophage A2]	89	64	321
572		2	1 571	sp P39237 Y05L_	HYPOTHETICAL 6.8 KD PROTEIN IN NRDC-TK INTERGENIC REGION.	89	47	570
646		914	1 459	gi 413982	ipa-58r gene product [Bacillus subtilis]	89	52	456
629	۳ 	1668	1 1901	gi 1107541	 C33D9.8 Caenorhabditis elegans	89 1	36	234
1 864	- 2	1510	1716	gi 145774	hsp70 protein (dnaK gene) [Escherichia coli]	89	48	207
920		098	1 432	gi 1510416	hypothetical protein (SP:P31466) [Methanococcus jannaschii]	89	54	429
952		1096	611	gi 603456	reductase [Leishmania major]	89 1	46	486
026		91	1 402	gi 1354775	pfos/R [Treponema pallidum]	89	46	312
1028		1064	534	gi 410117	diaminopimelate decarboxylase (Bacillus subtilis)	1 68	47	531
1029		428	1 216	gi 1335714 	Plasmodium falciparum mRNA for asparagine-rich antigen (clone 17Cl) [Plasmodium falciparum]	89	31	213
1058		692	348	gi 581649	epiC gene product [Staphylococcus epidermidis]	89	46	345
1 1096	7	1 665	1 465	gi 143434	Rho Factor (Bacillus subtilis)	89 1	43	201
1 1308	-	7	694	gi 1469939	group B oligopeptidase PepB [Streptococcus agalactiae]	1 68	50	693
1 1679		7	1 238	gi 517205	167 kDa Myosin-crossreactive streptococcal antigen [Streptococcus yogenes]	1 68	53	237
1 2039		m 	1 383	gi 153898	transport protein [Salmonella typhimurium]	89	51	381
1 2077		e -	326	pir C33496 C334	hisC homolog - Bacillus subtilis	89	47	324
2112	7	613	1 374	gi 64884	lamin LII [Xenopus laevis]	89	50	240
1 2273	-	1 793	398	gi 581648	epiB gene product [Staphylococcus epidermidis]	89	45	396
2948		2	1 385	gi 216869	Dranched-chain amino acid transport carrier [Pseudomonas aeruginosa] pir1A385341A38534 branched-chain amino acid transport protein braZ Pseudomonas aeruginosa	89	41	384
++		+	+			1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	+

Table?

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start Start (nt)	Stop (nt)	match acession	match gene name	sim -	* ident	length (nt)
1 2955		1 768	1 400	gi 904179	Inypothetical protein [Bacillus subtilis]	1 89	49	369
1 2981		572	1 288	gi 508979	GTP-binding protein (Bacillus subtilis)	- 89	48	285
3014	1	584	294	gi 1524394	ORF-2 upstream of gbsAB operon [Bacillus subtilis]	1 89	45	291
3082		336	169	gi 1204696	fructose-permease IIBC component [Haemophilus influenzae]	- 89	53	168
3108		103	1 258	gi 217855		1 89	48	156
1 3639	1	919	461	gi 1510490	Initrate transport permease protein [Methanococcus jannaschii]	1 89	47	459
1 3657	1	-	330	gi 155369	PTS enzyme-II fructose (Xanthomonas campestris)	1 89	48	330
3823		780	391	gi 603768 	HutI protein, imidazolone-5-propionate hydrolase [Bacillus subtilis] gi 603768 HutI protein, imidazolone-5-propionate hydrolase Bacillus subtilis	89	54	390
3982	1	2	1 277	gi 149435		1 89	47	276
4051	 		342	gi 450688 	hadM gene of Ecopriz gene product [Escherichia coli] pir 538437 538437 hadM protein - Escherichia coli pir 509629 509629 hypothetical protein A - Escherichia coli (SUB 40-520)	89	48	342
1 4089		12	1 209	gi 1353678	heavy-metal transporting P-type ATPase [Proteus mirabilis]	- 89	47	198
1 4143	-	1 47	1187	gi 603769	HutU protein, urocanase [Bacillus subtilis]	89	55	141
4148		2	352	gi 450688 	hsdM gene of Ecopril gene product [Escherichia coll] pir 538437 538437 hsdM protein - Escherichia coll pir S09629 S09629 hypothetical protein A - Escherichia coll (SUB 40-520)	89	51	351
4173		1 2	382	gi 1041097	Pyruvate Kinase (Bacillus psychrophilus)	89	48	381
4182		1 498	1 250	gi 413968	lipa-44d gene product (Bacillus subtilis)	- 89	50	249
4362		148	318	gi 450688 	hadM gene of EcoprrI gene product [Escherichia coli] pir S38437 S38437 hadM protein - Escherichia coli pir S09629 S09629 hypothetical protein A - Escherichia coli (SUB 40-520)	89	44	171
	111	1 9493	8300	gi 143727	[putative [Bacillus subtilis]	67	46	1194
1 31	111	110318	9833	gi 216746	D-lactate dehydrogenase [Lactobacillus plantarum]	67	41	486
1 32	m 	1560	3155	gi 1098557	renal sodium/dicarboxylate cotransporter [Homo sapiens]	1 69	46	1596
1 32	2	1 4945	1 4145	gi 1510720	[prephenate dehydratase [Methanococcus jannaschii]	67	51	801
36		1 5350	4268	gi 1146216 	145% identity with the product of the ORF6 gene from the Erwinia herbicola carotenoid biosynthesis cluster; putative (Bacillus subtilis)	67	28	1083
44	7	1 4492	1 5304	gi 1006621	hypothetical protein [Synechocystis sp.]	67	43	813

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

1, 1, 1, 1, 1, 1, 1,	Contig	ID ID	Start (nt)	Stop (nt)	match acession	match gene name	H E I	% ident	length (nt)
18 5002 4757 qiill030453 Profe Becillus subtilis 67			3943	8481		synthase large subunit precursor (Azospirillum brasilense) 22[846602 glutamate synthase (NADPH) (EC 1.4.1.13) alpha hain lum brasilense	67	52	4539
1 1.50 1.5	1 56	112	!	114678		TreR [Bacillus subtilis]	1 67	48	756
10 7570 6538 641854655 Na/H antiporter system [Bacillus alcalophilus] 67 67 67 67 67 67 67 6	62		1 5092	4757		[Bacillus, C-125, alkali-sensitive mutant 18224, Peptide Mutant,	67	45	336
1 11 11 11 11 11 11 11	1 62	110	17570	6338		Na/H antiporter system [Bacillus alcalophilus]	1 69 1	49	1233
19 5659 7176 91149432 purative [Lactilus subtilis] 67 11 11649 11039 91140847 [1899 game product [Bacillus subtilis] 67 11 11641 11514 91141396 11pa-52¢ gene product [Bacillus subtilis] 67 1 1 1164 91141304 91141304 918e-53d gene product [Bacillus subtilis] 67 1 1 1164 91141304 918e-1318 911526335 7mah [Bacillus subtilis] 67 2 4564 3559 91156681 Phypothetical protein (GB D26185_10) [Mycoplasama gentialium] 67 3 2899 2227 9114659 Phypothetical protein (GB D26185_10) [Mycoplasama gentialium] 67 4 5469 4198 911212775 GTP cyclohydrolase II (Bacillus amyloliquefaciens] 67 5 2913 2334 911303799 Yrkd [Bacillus subtilis] 67 6 1999 3728 911530379 Yrkd [Bacillus subtilis] 67 7 1720 1376 91144349 Phypothetical protein (Gphacia oberaceal) 67 8 6341 6673 91143139 Phys (Symethococcus sp.) 7 9 1351 1652 91141319 Phys (Symethococcus sp.) 7 1 1304 1418 91133347 Yrkd [Bacillus subtilis] 91144439 Phys (Symethococcus sp.) 9114444 Phys (Symethococcus sp.) 9114444 Phys (Symethococcus sp.) 9114444 Phys (Symethococcus sp.) 911444 Phys (Symethoc	66	- 3	2119	3321		(GB:GB:D90212_3)	1 69 1	20	1203
13 14549 14049 41 1408497	102	6	1 5695	1 7176		putative [Lactococcus lactis]	1 67	51	1482
15 14821 13592 Gil413976 Iiga-55r gene product [Bacillus subtilis] 67 67 67 67 67 67 67 6	103	113	!	114049		LP9D gene product [Bacillus subtilis]	1 67	48	501
17 14811 15194 911413993 15pa-5949 gene product [Bacillus subtilis] 67 1 1 1 1 1 1 1 1 1	109	115	1	113982		gene product (Bacillus	1 67	49	840
4 1713 2153 9411262335 Frank [Bacillus subtilis] 67 1 1 1 1149 (911143047 100RB [Bacillus subtilis] 67 1 1 1 1149 (911556885 Unknown (Bacillus subtilis] 67 1 1 1 1 1 1 1 1 1	109	117	114811	115194		gene product (Bacillus	67	29	384
1 1 1149 911143047 10RFB Bacillus subtilis]	121	4	1713	1 2153		YmaA [Bacillus subtilis]	1 67	54	441
5 4066 3518 gil556885 Unknown [Bacillus subtilis] 67 1 1 1 1 1 1 1 1 1	122		1	1 1149		ORFB [Bacillus subtilis]	1 67	35	1149
2 4584 3589 gill046081 hypothetical protein (GB:D26185_10) [Mycoplasma genitalium] 67 1 1 1 1 1 1 1 1 1	124	2	4060	3518		Unknown [Bacillus subtilis]	1 67	47	543
3 2899 2297 91 146549 kdpc [Escherichia coli] 67 4 5409 4198 91 121775	131	- 2	4584	1 3589			1 67	30	1 966
4 5409 4198 9111212775 GTP Cyclohydrolase II [Bacillus amyloliquefaciens] 67 8 6341 6673 91 1303709 YrkJ [Bacillus subtilis] 67 9 12720 3763 91 1377841	140	m -	1 2899	1 2297		[Escherichia	1 67 1	45	603
5 1913 1374 gill303709 YrkJ (Bacillus subtilis)	142	4	5409	4198		GTP cyclohydrolase II {Bacillus amyloliquefaciens}	1 67	55	1212
8 6341 6673 91 1377841	147	ر د	1 2913	2374		YrkJ (Bacillus subtilis]	1 67	44	540
4 12720 3763 gil 196319 SphX [Synechococcus sp.] 67	1 152	6 	6341	1 6673	gi 1377841	unknown [Bacillus subtilis]	1 67	48	333
6 1989 3428 gi 595681 12-oxoglutarate/malate translocator [Spinacia oleracea] 67 3 1351 1626 gi 1511101 shikimate 5-dehydrogenase [Methanococcus jannaschii] 67 2 917 1219 gi 142439 IATP-dependent nuclease [Bacillus subtilis] 67 110 112445 112801 ISP P37347 YECD HYPOTHETICAL 21.8 KD PROTEIN IN ASPS 5'REGION. 67 111 13047 114432 gi 732813	161	4	1 2720	1 3763		SphX (Synechococcus sp.]	1 69 1	47	1044
3 1351 1626 gill511101 Shikkmate 5-dehydrogenase [Methanococcus jannaschii] 67 2 917 2179 gill42439 ATP-dependent nuclease [Bacillus subtilis] 67 10 12445 12801 Sp P37347 YECD HYPOTHETICAL 21.8 KD PROTEIN IN ASPS 5'REGION. 67 11 13047 114432 gil 732813	163	9	1989	3428		translocator (Spinacia	1 69	47	1440
2 917 2179 ggil142439 IATP-dependent nuclease [Bacillus subtilis] 67	193		1351	1 1626		shikimate 5-dehydrogenase [Methanococcus jannaschii]	1 69	53	276
10 11445 12801 spp 93747 YECD	1 200	- 5	917	1 2179	gi 142439	[ATP-dependent nuclease [Bacillus subtilis]	1 69	48	1263
	1 206	110	112445	112801		21.8	67	47	357
2 1321 809 gill033037 1000 kDa heat shock protein (Hsp100) [Leishmania major] 67 3 1039 2052 gil809542 CbrB protein [Erwinia chrysanthemi] 67	1 206	111	113047	114432		[branched-chain amino acid carrier [Lactobacillus delbrueckii]	1 67	46	1386
3 1039 2052 gil809542	1 208	+ 2	1321	1 809	19111033037		1 67	36 1	513
	1 238		1039	1 2052		CbrB protein [Erwinia chrysanthemi]	1 67	42	1014 1

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

di I	I D	start (nt)	(nt)	match acession	match gene name	# S11	* ident	(nt)
1 246	1 2	176	367	gi 215098	excisionase [Bacteriophage 154a]	1 67	37	195
1 276	1 2	1 2260	1412	gi 303560	ORF271 (Escherichia coli	1 67	50	849
1 297	9	2223	3056	gi 142784		1 67	46	834
1 307	7	1 5220	4186	gi 1070013	protein-dependent [Bacillus subtilis]	19	43	1035
316	-	36	1028	gi 1161061	dioxygenase [Methylobacterium extorquens]	19	1 52	1 866
324	e	1 5650	1 5030	gi 1469784	[putative cell division protein ftsW [Enterococcus hirae]	1 67	49	621
1 336		524	1 264	gi 173122	urea amidolyase [Saccharomyces cerevisiae]	1 67	1 45	261
1 360		108	1394	splp30053 SYH_S	HISTIDYL-TRNA SYNTHETASE (EC 6.1.1.21) (HISTIDINETRNA LIGASE) (HISRS).	1 67	47	1287
364	e 	1 4890	3592	gi 151259 	HMG-CoA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir A44756 A44756 hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp.	67	46	1299
365	e 	2940	2113	gi 1296823	orf2 gene product [Lactobacillus helveticus]	1 67	47	828
1 367	7	325	918	git1039479	ORFU [Lactococcus lactis]	1 67	1 47	594
395	m -	999	1271	gi 1204516	hypothetical protein (GB:U00014_4) [Haemophilus influenzae]	1 67	5.5	1 909
415	1	1800	901	gi 882579	ICG Site No. 29739 [Escherichia coli]	1 67	46	006
419	1	1799	1 903	gi 520752	putative (Bacillus subtilis	19	48	897
474		7	962	gi 886906 	largininosuccinate synthetase [Streptomyces clavuligerus] pir S57659 S57659 argininosuccinate synthase (EC 6.3.4.5) - treptomyces clavuligerus	1 67	4 0	795
1 485	1 2	1921	2226	gi 143434	Rho Factor [Bacillus subtilis]	19	43	306
1 596	1	1728	1 865	gi 1303853	YqgF [Bacillus subtilis]	1 67	47	864
1 700	1	1 433	1 218	gi 1204628	hypothetical protein (SP:P21498) [Haemophilus influenzae]	19	1 47	216
908	1 2	249	647	gi 677947	AppC [Bacillus subtilis]	19	51	399
828	1 2	340	006	gi 777761	1rrA [Synechococcus sp.]	19	37	561
833	1	1407	916	gi 142996	regulatory protein [Bacillus subtilis]	19	41	492
856	1	1555	677	gil780224	[ZK970.2 [Caenorhabditis elegans]	1 67	38	777
888	1	1614	1 850	gi 437315	[TTG start codon [Bacillus licheniformis]	1 67	1 40	765
1034	1 1	1190	1 597	gi 1205113	hypothetical protein (GB:L19201 15) [Haemophilus influenzae]	1 67	45	594
1062	1 1	636	319	gi 1303850	YqgC [Bacillus subtilis]	1 67	41	318
1 1067	-	918	1 460	pir A32950 A329	r A32950 A329 probable reductase protein - Leishmania major	1 67	54	459

Table

300 507 204 1095 231 255 336 219 261 264 339 315 915 270 234 225 1269 1593 1251 969 1305 1032 774 | % ident | length (nt) 44 45 46 44 46 42 44 48 47 46 40 47 42 43 41 44 48 26 46 55 99 45 40 55 29 29 67 67 29 99 99 99 99 99 99 * sim 67 67 29 29 29 29 67 99 99 99 99 99 [67 kDa Myosin-crossreactive streptococcal antigen [Streptococcus yogenes] |HMG-CoA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir|A44756|A44756 molybdenum cofactor biosynthesis moeA protein [Methanococcus jannaschii] |ribonucleotide transport ATP-binding protein [Haemophilus influenzae] |deoxyribose-phosphate aldolase [Bacillus subtilis] pir|S49455|S49455 oligopeptide transport ATP-binding protein [Haemophilus influenzae] hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp. S. aureus - Putative coding regions of novel proteins similar to known proteins | deoxyribose-phosphate aldolase (EC 4.1.2.4) - acillus subtilis pir | S52915|S529 | nitrate reductase alpha chain - Bacillus subtilis (fragment) [hypothetical protein (SP:P31466) [Methanococcus jannaschii] |hypothetical protein (SP:P31805) [Haemophilus influenzae] |mercury scavenger protein [Haemophilus influenzae] |uroporphyrinogen III methyltransferase [Zea mays] (cystathionine beta-lyase [Arabidopsis thaliana] [inducible nitric oxide synthase [Gallus gallus] |cystathionine beta-lyase [Arabidopsis thaliana] [peptide-synthetase [Amycolatopsis mediterranei] |ATP-dependent nuclease [Bacillus subtilis] [hypothetical protein [Synechocystis sp.] hypothetical protein [Bacillus subtilis] [phaD gene product [Rhizobium meliloti] GTG start codon [Lactococcus lactis] |SmtB [Synechococcus PCC7942] |ORF 1 [Mycoplasma mycoides] [menD [Bacillus subtilis] |Codv [Bacillus subtilis] [mucin [Homo sapiens] match gene name acession fgi | 1001369 |gi|1510416 |gi|1204545 |gi|1072398 | 5 | 3810 | 3265 |gi|1204323 |gi|1205366 gi|1531541 |gi|1483199 [gi|1510751 |gi|1205337 |gi|557489 |gi|517205 |gi|308861 |gi|151259 |gi|704397 |gi|535348 |gi|292046 1911150209 |gi|665999 1gi | 809660 | 1 | 1 | 1305 | gi|142440 1gi1998342 | 9236 | 8205 |gi|704397 |gi|46491 110124 1 7178 1 8428 9258 1 2868 | 5143 | 4370 114 (111693 (10998 3989 1 2542 1 7826 6347 | 5253 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 302 1 265 340 396 507 234 256 393 251 398 1276 1 3 | 3075 9034 9888 1 2273 1 8059 454 464 1 2 | 584 658 1 1 82 <u>ო</u> . -1 1 728 3000 | 1 | 1 - 2 1 1 | 2 ~ -_ -7 9 6 110 9 œ — 9 2181 | 1 3066 | 1 3101 | 1 113 3087 3598 3788 3765 3926 1358 3883 4417 15 31 31 32 44 48 28 62 9/ 91 ~

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

1	1 1 1 1 1	1 1 1 1 1 1 1 1		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				+
Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	e sim	% ident	length (nt)
103	4	3418	2732	gi 971344 	nitrate reductase gamma subunit (Bacillus subtilis) sp P42177 NARI BACSU NITRATE REDUCTASE GAMMA CHAIN (EC 1.7.99.4). gi 1009369 Respiratory nitrate reductase (Bacillus subtilis) (SUB -160)	99	48	687
109	φ 	4243	4674	gi 170886 	glucosamine-6-phosphate deaminase (Candida albicans) pir[A46652[A46652] glucosamine-6-phosphate isomerase (EC 5.3.1.10) - east (Candida albicans)	99	45	432
112	117	117491	117712	gi 1323179	ORF YGR111w [Saccharomyces cerevisiae]	99	33	222
116	2	1 4667	1 2637	gi 1491813	gamma-glutamyltranspeptidase (Bacillus subtilis)	99	43	2031
150	5	3189	1 2989	19111146224	putative [Bacillus subtilis]	99	30	201
172	2	3264	1 3662	911755152	highly hydrophobic integral membrane protein [Bacillus subtilis] sp 842953 TAGG_BACSU TEICHOIC ACID TRANSLOCATION PERWEASE PROTEIN AGG.	99	41	399
174	5 -	4592	3723	gi 1146241	pantothenate synthetase [Bacillus subtilis]	99	49	870
175	4	3209	1 2880	gi 642655	unknown [Rhizobium meliloti]	99	29	330
175	111	8743	1 7994	gi 854655	Na/H antiporter system [Bacillus alcalophilus]	99	43	750
190		9707	5727	gi 451072	di-tripeptide transporter [Lactococcus lactis]	99	40	1353
195	115	13919	113713	gi 1322411	unknown [Mycobacterium tuberculosis]	99	42	207
217	. 3	2822	1 2595	gi 1143542	alternative stop codon [Rattus norvegicus]	99	36	228
233	6	7133	6135	gi 1458327	F08F3.4 gene product (Caenorhabditis elegans)	99	47	1 666
1 238	-	43	1041	gi 809541	(CbrA protein (Erwinia chrysanthemi)	99	42	666
1 241	-	2102	1 1053	gi 153067	peptidoglycan hydrolase [Staphylococcus aureus]	99	53	1050
261	-	1178	648	gi 1510859	M. jannaschii predicted coding region MJ0790 [Methanococcus jannaschii]	99	40	531
263	3	3731	1 2973	gi 1205865	tetrahydrodipicolinate N-succinyltransferase (Haemophilus influenzae)	99	47	759
272	œ 	6548	5484	gi 882101 	high affinity nickel transporter (Alcaligenes eutrophus) sp P23516 HOXN_ALCEU HIGH-AFFINITY NICKEL TRANSPORT PROTEIN.	99	44	1065
1 276	3	1 2805	1 2104	gi 1208965	hypothetical 23.3 kd protein [Escherichia coli]	99	47	702
278	- 2	1 2830	1 1784	gi 1488662	phosphatase-associated protein [Bacillus subtilis]	99	48	1047
1 278		3830	1 2952	gi 303560	ORF271 [Escherichia coli]	99	45	879
279	5	3894	1 2218	gi 1185289 	2-succiny1-6-hydroxy-2,4-cyclohexadiene-1- carboxylate synthase [Bacillus subtilis]	99	48	1677
1 288	4	1 2535	1 2275	gi 1256625	putative [Bacillus subtilis]	99	42	261
1 292	2	1133	942	gi 1511604	M. jannaschii predicted coding region MJ1651 (Methanococcus jannaschii)	99	30	192
								-

Table ?

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig ID	ORF	Start (nt)	Stop (nt)	match acession	match gene name	ae − .	TOGUE	(nt)
294		1116	1 559	gi 216314	esterase (Bacillus stearothermophilus]	99	45	258
297	4	2913	1978	gi 994794 	Cytochrome a assembly facto [Bacillus subtilis] sp P24009 COXX_BACSU PROBABLE CYTOCHROME C OXIDASE ASSEMBLY FACTOR.	99	45	936
316	4	1 2053	1 2682	gi 1107839	alginate lyase [Pseudomonas aeruginosa]	99	1 40	630
338	4	1 2460	1 2302	gi 520750	biotin synthetase (Bacillus sphaericus)	99	28	159
339		1214	1 735	gi 467468	7, 8-dihydro-6-hydroxymethylpterin-pyrophosphokinase [Bacillus ubtilis]	99	52	480
363			1 863	gi 581649	lepiC gene product [Staphylococcus epidermidis]	99	1 47	861
366	2	1 232	1 483	gi 1103505	unknown [Schizosaccharomyces pombe]	99	53	252
367	4	1 2468	1845	sp P20692 TYRA_	PREPHENATE DEHYDROGENASE (EC 1.3.1.12) (PDH).	99	1 80	624
372	8 -	1 2150	1599	gi 467416	unknown [Bacillus subtilis]	99	38	552
378		1 212	1009	gi 147309	purine nucleoside phosphorylase [Escherichia coli]	99	20	198
401	·		1 462	gi 388263 	p-aminobenzoic acid synthase [Streptomyces griseus] pir JN0531 JN0531 p- aminobenzoic acid synthase - Streptomyces riseus	99	46	462
404	7	1 4826	5254	gi 606744	cytidine deaminase [Bacillus subtilis]	99	51	429
411	1 2	1738	1103	gi 1460081	unknown [Mycobacterium tuberculosis]	99	1 44	969
420		7	541	gi 1046024	Na+ ATPase subunit J [Mycoplasma genitalium]	99	1 49	1 540
431	-		858	gi 1500008	[M. jannaschii predicted coding region MJ1154 [Methanococcus jannaschii]	99	1 30	828
443	7 1	1 5679	1 5299	gi 852076	MrgA [Bacillus subtilis]	99	1 46	381
444	m 	3405	2413	gi 153047 	1ysostaphin (ttg start codon) Staphylococcus simulans pir A25881 A25881 1ysostaphin precursor - Staphylococcus simulans sp P10547 LSTP_STASI LYSOSTAPHIN PRECURSOR (EC 3.5.1).	99	51	666
561		926	1 480	(gi/1204905	DNA-3-methyladenine glycosidase I [Haemophilus influenzae]	99	45	477
562	m 	1066	1383	gi 1046082	IM. genitalium predicted coding region MG372 [Mycoplasma genitalium]	99	52	318
576		11	1 724	gi 305014	IORF_0234 [Escherichia coli]	99	43	714
577	۳ -	1190	1 903	gi 1001353	hypothetical protein [Synechocystis sp.]	99	52	288
584		2	331	sp P24204 YEBA_	HYPOTHETICAL 46.7 KD PROTEIN IN MSBB-RUVB INTERGENIC REGION (ORFU).	99	48	330
592		1410	1 706	gi 928839	ORF266; putative [Lactococcus lactis phage BK5-T]	99	51	705
601	-	1433	1 720	qi 1488695	Inovel antigen: orf-2 [Stabhylococcus aureus]		r.	714

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	e sis	% ident	length (nt)
619	<u>ه</u> ا	468	1 845		similar to M. musculus transport system membrane protein, Nramp PIR:A40739) and S. cerevisiae SMF1 protein (PIR:A45154) Caenorhabditis elegans]	99	45	378
706	- 5	561	355	gi 804808	unknown protein (Rattus norvegicus)	99	46	207
734	5	1 673	512		phosphatidylcholine binding immunoglobulin heavy chain IgM variable region [Mus musculus]	99	09	162
740	 	3	317	gil1209272	argininosuccinate lyase [Campylobacter jejuni]	99	42	315
764		310	1 747	gi 435296 	alkaline phosphatase like protein [Lactococcus lactis] pir 839339 839339 alkaline phosphatase-like protein - Lactococcus actis	99	42	438
852		1 338	171	1911536955	CG Site No. 361 [Escherichia coli]	99	43	168
988		8	158	gi 289272	ferrichrome-binding protein [Bacillus subtilis]	99	44	156
688		1 462	1 232	gi 833061	HCMVUL77 (AA 1-642) [Human cytomegalovirus]	99	99	231
893		2	1 247	gi 149008	putative [Helicobacter pylori]	99	45	246
006	1 - 1	1425	1 733	gi 580842	F3 [Bacillus subtilis]	99	51	693
906	7	1 2300	1473	gi 790945	aryl-alcohol dehydrogenase [Bacillus subtilis]	99	53	828
947		67	549	gi 410117		99	47	471
950		1100	552	gi 48713	orf145 [Staphylococcus aureus]	99	35	549
955	5	68	475	gi 1204390	uridine kinase (uridine monophosphokinase) [Haemophilus influenzae]	99	50	387
981	- 5	1308	1 997	gi 457146	rhoptry protein [Plasmodium yoelii]	99	38	312
986	-	1 25	315	gi 305002	ORF_f356 [Escherichia coli]	99	31	291
1057		e -	203	gi 1303853	[YqgF (Bacillus subtilis]	99	40	201
1087	- 1		1 294	gi 575913	unknown [Saccharomyces cerevisiae]	99	53	294
1105			231	gi 1045799	[methylgalactoside permease ATP-binding protein [Mycoplasma genitalium]	99	46	231
1128		2	1 574	gi 1001493	hypothetical protein [Synechocystis sp.]	99	46	573
1150	1	498	1 250	gi 1499034	M. jannaschii predicted coding region MJ0255 [Methanococcus jannaschii]	99	40	249
1180	2	1 707	453	gi 215908	DNA polymerase (g43) [Bacteriophage T4]	99	46	255
1208		1123	1 587	gi 1256653	[DNA-binding protein [Bacillus subtilis]	99	58	537
1342			1 402	gi 1208474	hypothetical protein [Synechocystis sp.]	99	53	402
1761	2 -	- 589	1 398	gi 215811	tail fiber protein [Bacteriophage T3]	99	50	192

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

1	1			1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		+	+	1 1 1 1 1 1 1
Contig	ORF IID	Start (nt)	Stop (nt)	match	match gene name	8 sim	% ident	length (nt)
1983	1	499	1 251	gi 1045935	DNA helicase II [Mycoplasma genitalium]	99	40 1	249
1 2103	- 2	176	1 400	gi 929798		99	46	225
1 2341	. — .	373	188	gi 1256623	[exodeoxyribonuclease [Bacillus subtilis]	99	38	186
1 2458		325	1 164	gi 1019410	unknown [Schizosaccharomyces pombe]	99	47	162
1 2505		468	1 235	gi 1510394		99	39	234
1 2525	-	558	1 280	gi 1000695	loytotoxin L [Clostridium sordellii]	1 99	44	279
2935		8	1 275	gi 765073	autolysin [Staphylococcus aureus]	99	47	273
3005		114	1 305	gi 1205784	heterocyst maturation protein [Haemophilus influenzae]	99	46	192
3048		80	772	gi 1303813	YqeW (Bacillus subtilis)	99	42	198
3071		1	189	gi 1070014	protein-dependent (Bacillus subtilis]	99	41	189
3081	-	404	1 225	gi 984212	unknown [Schizosaccharomyces pombe]	99	44	180
1 3090	7	280	1 386	gi 1204987	DNA polymerase III, alpha chain [Haemophilus influenzae]	99	48	195
3318		-	1 387	lgi11009366	Respiratory nitrate reductase [Bacillus subtilis]	99	49	387
3739		1 798	1 400	gi 1109684	Prov (Bacillus subtilis)	99	47	399
3796		1 402	1 202	gi 853760	acyl-CoA dehydrogenase [Bacillus subtilis]	1 99	1 09	201
3924		595	1 347	gi 563952	[gluconate permease [Bacillus licheniformis]	99	46	249
4240		e 	350	gi 151259 	HMG-COA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir A44756 A44756 hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp.	99	51	348
4604		7	1 234	ріг А26713 ВННС	hemocyanin subunit II - Atlantic horseshoe crab	99	46	228
4	6	8845	9750	 gi 145646	cynR [Escherichia coli]	65	35	1 906
9	2	2708	1 3565	lgi1887824	ORF_0310 [Escherichia coli]	65	47	828
13	-	1993	866	gi 143402 	recombination protein (ttg start codon) [Bacillus subtilis] gi 1303923 RecN [Bacillus subtilis]	65	44	1 966
1 15	7	2493	3524	gi 1403126	czcD gene product (Alcaligenes eutrophus)	65	38	1032
18	. —	1908	1 1372	gi 349187	acyltransferase [Saccharomyces cerevisiae]	65	50	537
21	m ———	1467	2492	gi 149518 	phosphoribosyl anthranilate transferase (Lactococcus lactis) pir[335126 335126 anthranilate phosphoribosyltransferase (EC .4.2.18) - Lactococcus Lactis subsp. Lactis	65	52	1026
1 25	4	3374	4312	gi 1502420	malonyl-CoA:Acyl carrier protein transacylase [Bacillus subtilis]	65	44	939
	1 1 1 1	1				+	+	+

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORE	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
1 27		1 390	626	gi 1212729	YqhJ [Bacillus subtilis]	65	45	237
31	112	111040	110387	gi 509245	D-hydroxyisocaproate dehydrogenase [Lactobacillus delbrueckii]	65	41	654
38	124	119172	119528	gi 547519	H-protein [Flaveria cronquistii]	65	41	357
44	7	1 790	1746	gi 405882		65	46	957
44	112	9356	8832	gi 1205905	molybdenum cofactor biosynthesis protein [Haemophilus influenzae]	65 1	50	525
1 45	ω	1 6635	1 7588	lgi 493074	ApbA protein [Salmonella typhimurium]	65	46	954
51	7	280	1503	gi 580897	OppB gene product [Bacillus subtilis]	65	45	924
1 52		1 225	953	gi 1205518	NAD(P)H-flavin oxidoreductase [Haemophilus influenzae]	65	45	729
- 55	4	1339	1058	pir A44459 A444		65	41	282
1 67	6	1 7421	8272	gi 143607	Isporulation protein [Bacillus subtilis]	65	42	852
7.3		4446	1 5375	gi 1204896	1 1 1 1 1 1 1 1 1	65	37	930
1 74	-	954	1 478	gi 1204844	H. influenzae predicted coding region HI0594 [Haemophilus influenzae]	65	20	477
77		7	1757	gi 1046082	[M. genitalium predicted coding region MG372 [Mycoplasma genitalium]	65	46	756
7.7	2,	1 795	1433	gi 1222116	permease [Haemophilus influenzae]	65	37	639
81	. . .	1 4728	3454	gi 1001708	hypothetical protein [Synechocystis sp.]	65	49	1275
91	7	8548	8357	lgi11399263	cystathionine beta-lyase [Emericella nidulans]	65	40	192
86	. —	1608	1 1988	gi 467423	unknown [Bacillus subtilis]	65	38	381
86	4	1 2250	1 2987	gi 467424	unknown [Bacillus subtilis]	65	45	738
102	e 	1 2598	2119	gi 1511532 	N-terminal acetyltransferase complex, subunit ARD1 [Methanococcus jannaschii]	65	39	480
102	4	3647	1 2862	gi 1204637	H. influenzae predicted coding region HI0388 [Haemophilus influenzae]	65	32	786
103	9	10851	9841	gi 142695 	S-adenosyl-L-methionine:uroporphyrinogen III methyltransferase Bacillus megaterium]	65	47	1011
103	110	10439	110119	gi 710021	nitrite reductase (nirD) [Bacillus subtilis]	65	51	321
106	7	1 262	11140	gi 39881	ORF 311 (AA 1-311) (Bacillus subtilis)	65	44	879
109		1 3909	1 4268	gi 1204399	glucosamine-6-phosphate deaminase protein [Haemophilus influenzae]	65	44	360
109	01-	1 7165	1 8595	gi 536955	[CG Site No. 361 [Escherichia coli]	65	41	1431
		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1						

Table 2

414 150 579 762 420 1926 540 1911 2637 2496 1296 630 1515 1383 228 201 252 642 984 198 612 774 339 987 | % ident | length (nt) 20 99 39 46 41 40 46 40 45 25 32 26 27 37 20 46 49 38 53 41 40 46 20 43 65 65 65 65 65 65 9 65 65 65 65 65 65 65 65 65 65 65 65 65 65 65 65 65 l & sim |beta-1,4-N-acetylmuramoylhydrolase [Enterococcus hirae] pir|A42296|A42296 POLY (GLYCEROL-PHOSPHATE) LPHA-GLUCOSYLTRANSFERASE (EC 2.4.1.52) (TECHOIC |rodD (gtaA) polypeptide (AA 1-673) [Bacillus subtilis] pir|S06048|S06048 | probable rodD protein - Bacillus subtilis sp|P13484|TAGE_BACSU FROBABLE IH. influenzae predicted coding region HI0318 [Haemophilus influenzae] |stringent response-like protein [Streptococcus equisimilis] | pir|S39975|S39975 stringent response-like protein - Streptococcus | lysozyme 2 (EC 3.2.1.-) precursor - Enterococcus irae (ATCC 9790) ribonuclease HII (EC 31264) (RNASE HII) [Haemophilus influenzae] |DNA or RNA helicase, DNA-dependent ATPase [Bacillus subtilis] |hypothetical protein (GB:U00022_9) (Haemophilus influenzae] |mutator mutT (AT-GC transversion) [Escherichia coli] transcriptional activator protein (Bacillus brevis) [homologous to sp:HTRA_ECOLI [Bacillus subtilis] |repeat organellar protein (Plasmodium chabaudi] ornithine acetyltransferase (Bacillus subtilis) exonuclease V alpha-subunit [Escherichia coli] [5'guanylate kinase [Haemophilus influenzae] [alanyl-tRNA synthetase [Escherichia coli] ORF IV (AA 1-489) [Figwort mosaic virus] |ORF3; putative [Rhodobacter capsulatus] |D02_orf569 [Mycoplasma pneumoniae] |ORF1 [Streptococcus equisimilis] ACID BIOSYNTHESIS ROTEIN E). |putative [Bacillus subtilis] |Orf2 [Streptomyces griseus] [HisBd [Lactococcus lactis] [precursor [Homo sapiens] [fecD [Escherichia coli] match gene name quisimilis acession |gi|1139574 |gi|1064809 |gi|1209768 |gi|1146225 |gi|1146200 |gi|1204989 |qi||1151158 |gi|1205308 |gi|1204571 |gi|1205974 match 3915 |gi|407881 |gi|407880 |gi|710496 |gi|216513 |gi|151943 |gi|145220 |gi|408115 |gi|580920 1gi|149379 |gi|145927 1gi1148304 | 8104 |gi|882711 |gi|37589 |gi|58812 2841 5272 118191 1 4295 3215 3751 3794 4231 | 4380 1 8640 7004 1 3587 | 3838 1948 4785 1 2327 | 3709 2513 Start | Stop 111288 112658 1 4178 1 4195 (nt) 1 940 422 1 202 3688 3195 (nt) 3482 3844 5265 1 4780 3882 9218 112049 111 |10733 7543 3858 1 4982 1 7908 110599 1902 1 4968 1 1 1 1278 116896 - 5 m Contig |ORF ID |ID -œ — 5 4 - 5 9 _ 4 9 თ 9 116 2 2 <u>ش</u> 4 <u>۔</u> ص 110 112 145 195 110 110 110 112 112 125 126 127 143 150 166 188 189 195 206 217 236 237 241

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

111111	1	1			A	+	+	+
Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	e sin	% ident	length (nt)
1 274		3	278	lgi1496558	orfX [Bacillus subtilis]	65	42	276
301	- 2	1 982	815	gi 467418	unknown [Bacillus subtilis]	65	45	168
1 307	4	3586	1 2864	gi 1070014		65	40	723
335	N 	2286	1399	gi 146913	N-acetylglucosamine transport protein [Escherichia coli] pir B2995 WQEC2N phosphotransferase system enzyme II (EC .7.1.69), N-acetylglucosamine- specific - Escherichia coli sp P09323 PTAA_ECOLI PTS SYSTEM, N- ACETYLGLUCOSAMINE-SPECIFIC ITABC OMPONENT (EILA	65	50	888
1 338	S -	4120	3170	gi11277029	biotin synthase [Bacillus subtilis]	65	49	951
1 343		1490	1 2800	gi 143264	 membrane-associated protein (Bacillus subtilis)	65	48	1311
344	4	1 2761	2531	gil1050540		65	34	231
1 358	m —	3421	3621	gi 1146220	NAD+ dependent glycerol-3-phosphate dehydrogenase [Bacillus subtilis]	65	47	201
1 364	1	1 238	669	gi 1340128	ORF1 [Staphylococcus aureus]	65	51	462
379	 		576	gi 143331	alkaline phosphatase regulatory protein (Bacillus subtilis)	65	0	576
1 379	- 3	3666	4346	gi 143268	dihydrolipoamide transsuccinylase (odhB; EC 2.3.1.61) (Bacillus ubtilis)	65	20	681
1 428		187	483	gi 1420465	ORF YOR195w [Saccharomyces cerevisiae]	65	45	297
1 438	7 7	1 272	838	gi 143498	degS protein [Bacillus subtilis]	65	38	567
1 444	17.	1 9280	110215	gi 1204756	ribokinase [Haemophilus influenzae]	65	47	936
449	7	1241	1531	gi 599848		65	41	291
478	2	1452	1 865	gi 1045942	glycyl-tRNA synthetase [Mycoplasma genitalium]	65	39	588
1 479		1 1032	1 517	gi 1498192		65	40	516
480	9	4312	1 5637	gi 415662 	UDP-N-acetylglucosamine 1-carboxyvinyl transferase (Acinetobacter alcoaceticus)	65	84	1326
1 484		5	1 430	gi 146551	(transmembrane protein (kdpD) [Escherichia coli]	65	44	429
1 499		1 54	932	gi 603456	reductase [Leishmania major]	65	53	879
1 505		914	1 459	gi 1518853	OafA [Salmonella typhimurium]	65	39	456
571	5	1509	883	gi 49399 	open reading frame upstream glnE [Escherichia coli] ir 537754 537754 hypothetical protein XE (glnE 5' region) - cherichia coli	65	44	627
611	1 2	1 506	1 270	gi 10961		65	40	237

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start	Stop	-+	match dene name	sim s		length /
O.	QI.	(nt)	(nt)	acession		,		(nt)
705	-	564	1 283	gi 710020	Initrite reductase (nirB) [Bacillus subtilis]	65	52	282
712			177	gi 289272	[ferrichrome-binding protein [Bacillus subtilis]	65	37	177
712	- 5	196	1 354	gi 289272	[ferrichrome-binding protein [Bacillus subtilis]	9	37	159
743	г г - — -	7	631	gi 310631	ATP binding protein [Streptococcus gordonii]	69	45	630
749	2	393	677	lgi1467374	single strand DNA binding protein [Bacillus subtilis]	65	29	387
762		1698	1 850	gi 160399	multidrug resistance protein [Plasmodium falciparum]	65	48	849
788		85	315	gi 1129096	lunknown protein (Bacillus sp.)	65	35	231
850		F-1	408	gi 1006604	hypothetical protein (Synechocystis sp.)	65	37	408
806			444	gi 1199546		65	46	444
925		-	174	gi 1256653	[DNA-binding protein [Bacillus subtilis]	65	54	174
1031		26	232	gi 238657	<pre> AppC=cytochrome d oxidase, subunit I homolog [Escherichia coli, K12, eptide, 514 aa]</pre>	65	47	207
1037		414	1 262	gi 1491813	gamma-glutamyltranspeptidase [Bacillus subtilis]	65	46	153
1053	. —	348	175	gi 642655	unknown [Rhizobium meliloti]	65	34	174
1149		1399	1 752	gi 1162980	ribulose-5-phosphate 3-epimerase [Spinacia oleracea]	99	48	648
1214		881	1 495	gi 1205959	lactam utilization protein [Haemophilus influenzae]	65	45	387
1276		1 476	276	pir 835493 8354	site-specific DNA-methyltransferase StsI (EC 2.1.1) - Streptococcus sanguis	65	35	201
1276	7	006	577	gi 473794	'ORF' [Escherichia coli]	65	34	324
2057		272	138	1911633699	TrsH [Yersinia enterocolitica]	65	21	135
2521		336	169	gi 1045789	hypothetical protein (GB:U14003_76) [Mycoplasma genitalium]	65	41	168
2974		1 590	1 297	gi 152052	enantiomerase-selective amidase [Rhodococcus sp.]	65	45	294
3031		306	154	pir JQ1024 JQ10	hypothetical 30K protein (DmRP140 5' region) - fruit fly (Drosophila melanogaster)	65	45	153
3069		m 	278	gi 144906 	product homologous to E.coli thioredoxin reductase: J.Biol.Chem. 1988) 263:9015-9019, and to F52a protein of alkyl hydroperoxide eductase from S.typhimurium: J.Biol.Chem. (1990) 265:10535-10540; pen reading frame A	65	46	276
3146		282	142	gi 49315	ORF1 gene product [Bacillus subtilis]	65	47	141
3170		629	341	gi 1507711	indolepyruvate decarboxylase [Erwinia herbicola]	65	44	339
	++	++	+		,		+	

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF I	Start (nt)	Stop	match acession	match gene name		* ident	length (nt)
3546			1 303	-+	hsdM gene of EcoprII gene product [Escherichia coli] pir S38437 S38437 hsdM protein - Escherichia coli pir S09629 S09629 hypothetical protein A - Escherichia coli (SUB 40-520)	65	42	303
3782	-	2	328	gi 166412	NADH-glutamate synthase [Medicago sativa]	65	42	327
1 3990	-	374	189	 gi 1009366	Respiratory nitrate reductase [Bacillus subtilis]	65	53	186
1 4032	-	613	308	gi 1323127	ORF YGR087c [Saccharomyces cerevisiae]	65 1	20	306
1 4278	2	726	364	1911197667	vitellogenin (Anolis pulchellus)	65	42	363
1 19	4	4259	5518	gi 145727	deaD [Escherichia coli]	1 69	45	1260
1 19	9	7639	1 6926	gi 1016232	ycf27 gene product [Cyanophora paradoxa]	1 69	36	714
1 20	8	1 7053	6454	gi 765073	autolysin [Staphylococcus aureus]	64	4 7 4	009
31	113	112706	111537	gi 414009	ipa-85d gene product [Bacillus subtilis]	64	45 +	1170
1 33	7	2388	4364	gi 1204696	fructose-permease IIBC component [Haemophilus influenzae]	1 69	47	1 7761
1 36	8 -	1871	3013		glutamate permease [Escherichia coli]	64	40 4	1143
1 37	9	4065	4409	gi 39815	orf 2 gene product [Bacillus subtilis]	1 69	46	345
1 45	6	7852	1 8760	gi 1230585	Inucleotide sugar epimerase [Vibrio cholerae 0139]	64	53	1 606
53	8	1540	1 1899	gi 1303961	الالإلام (Bacillus subtilis)	64 +	20	360
1 56	9	4793	3855	gi 457514	gltC [Bacillus subtilis]	64	45	626
1 56	124	130002	130247	gi 470331	similar to zinc fingers [Caenorhabditis elegans]	1 69	42	246
1 62	4	2759	1 2421	gi 642655	unknown [Rhizobium meliloti]	64	28	339
85	9	1 7178	6027	gi 457702 	5-aminoimidazole ribonucleotide-carboxilase [Pichia methanolica] pir 539112 539112 phosphoribosylaminoimidazole carboxylase (EC .1.1.21) - yeast (Pichia methanolica)	64	46	1152
96	6	9251	110030	gi 1511513	ABC transporter, probable ATP-binding subunit (Methanococcus jannaschii)	64	42	780
100	1	1	009	gi 765073	autolysin [Staphylococcus aureus]	64	44	009
106	1.5	3868	4854	gi 466778	lysine specific permease [Escherichia coli]	64	46	1 286
123	2	838	554	gi 467484	unknown [Bacillus subtilis]	64	47	285
127	8	7514	7810	gi 210061 	serotype-specific antigen [African horse sickness virus] pir S27891 S27891 capsid protein VP2 - African horse sickness virus	64	28	297
131	7	7134	6721	gi 1511160	[M. jannaschii predicted coding region MJ1163 [Methanococcus jannaschii]	64	46	414

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contia	+	Start	Stop	-+	match oene name	++	* ident 1	length
	01	(nt)	(nt)	acession			- +	(nt)
1 142	5	5455	1 4817	gi11173517	riboflavin synthase alpha subunit [Actinobacillus pleuropneumoniae]	64	44	639
143	-	109	1 356	pir A32950 A329	probable reductase protein - Leishmania major	64	52	354
149	110	3555	1 3295	gi 398151	major surface antigen MSG2 [Pneumocystis carinii]	64	44	261
1 154	4	3134	1 2307	gi 984587	Dinp {Escherichia coli}	64	50	828
1 161	2	3855	1 4880	gi 903304	ORF72 (Bacillus subtilis]	64	37	1026
165		33	1 791	gi 467483	unknown [Bacillus subtilis]	1 64	38	759
175	9	6355	1 4844	gi 1072398	phaD gene product [Rhizobium meliloti]	64	42	1512
188	۳ -	1 2042	1 2500	gi 1001961	MMC class II analog (Staphylococcus aureus)	1 64	45	459
1 195	114	13667	13446	gi 396380	No definition line found [Escherichia coli]	64	47	222
1 206	115	116429	116938	gi 304134	largC [Bacillus stearothermophilus]	64	49	510
1 215		1 560	1 282	gi 142359	ORF 6 (Azotobacter vinelandii)	64	39	279
1 243	7	1 7818	6928	gi 414014	lipa-90d gene product [Bacillus subtilis]	64	49	891
1 258	7	1330	1 845	gi 664754	P17 [Listeria monocytogenes]	64	38	486
1 259	1	1 462	232	gi 1499663	[M. jannaschii predicted coding region MJ0837 [Methanococcus jannaschii]	1 64 1	52	231
1 263	9	6565	1 5567	gi 142828 	laspartate semialdehyde dehydrogenase [Bacillus subtills] sp Q04797 DHAS_BACSU ASPARTATE-SEMIALDEHYDE DEHYDROGENASE (EC .2.1.11) (ASA DEHYDROGENASE).	64	48	666
271		3	1163	gi 467091	hflX; B2235_C2_202 [Mycobacterium leprae]	64	44	1161
280		173	1450	gi 1303839	YqfR [Bacillus subtilis]	64	43	1278
293	1	1 2532	1 1267	gi 147345	primosomal protein n' [Escherichia coli]	64	45	1266
295	2	742	1488	gi 459266	Potential membrane spanning protein (Staphylococcus hominis) pir15429321542932 potential membrane spanning protein - taphylococcus hominis	64	39	747
301	2	1625	1446	gi 580835	lysine decarboxylase (Bacillus subtilis]	64	35	180
315	1 4	5064	3949	gi 143396	quinol oxidase (Bacillus subtilis]	64	45	1116
321	1	1264	635	gi 710496	transcriptional activator protein [Bacillus brevis]	64	41	630
333	2	4520	1 4239	gi 1314295	ORF2; putative 19 kDa protein [Listeria monocytogenes]	1 64 1	43	282
342	1	. – .	1 549	gi 142940	[ftsA [Bacillus subtilis]	64	38	549
353	<u>م</u>	1 2878	1 2324	gi 537049	ORF_0470 [Escherichia coli]	64	44	555
							•	

Table ?

| % sim | % ident | length (nt) |cyclomaltodextrin glucanotransferase [Bacillus stearothermophilus] i|39835 phosphotransferase system glucose-specific enzyme II (Bacillus subtilis) (pir|S25295|A328 |oxoglutarate dehydrogenase (lipoamide) (EC 1.2.4.2) - Bacillus subtilis lenterotoxin type E precursor (Staphylococcus aureus) pir/A28179|A28179|
l enterotoxin E precursor - Staphylococcus aureus sp/P12993|ETXE_STAAU | ENTEROTOXIN TYPE E PRECURSOR (SEE). 1H. influenzae predicted coding region HI0238 [Haemophilus influenzae] spermidine/putrescine transport system permease protein [Haemophilus |fibrinogen-binding protein [Staphylococcus aureus] pir|S34270|S34270 |dicarboxylic amino acids Dip5p permease [Saccharomyces cerevisiae] cyclomaltodextrin glucanotransferase (Bacillus earothermophilus) |bacterioferritin comigratory protein [Haemophilus influenzae] |pir|A36933|A369 |diacylglycerol kinase homolog - Streptococcus mutans [purine nucleoside phosphorylase [Escherichia coli] fibrinogen-binding protein - Staphylococcus ureus |NADH dehydrogenase F [Streptogyna americana] |host interacting protein (Bacteriophage B1) |arginyl tRNA synthetase [Bacillus subtilis] |hypothetical protein [Synechocystis sp.] [HhdA precursor [Haemophilus ducreyi] [ahrC protein [Bacillus subtilis] |DNA primase [Bacillus subtilis] |ORF o162 [Escherichia coli] |pfoS/R [Treponema pallidum] |Rap60 [Bacillus subtilis] |OrfX [Bacillus subtilis] |sp|P36929|FMU E |FMU PROTEIN. | match gene | influenzae] acession |gi|1369948 |gi|1205582 |gi|1204496 gi|1204511 |gi|1213234 |gi|1001383 |gi|1049115 |gi|1151072 |gi|1146177 |gi|1354775 1gi | 142450 |gi|311976 gi1969026 1gi1147309 1gi1606376 gil755823 |gi|142865 |gi|971336 |gi|153002 |gi|39833 (nt) | 591 1 917 1 963 Start 1 1 1 2217 (nt) 1 2 | 1172 1 2020 1 4082 1 1035 4 | 1911 4 | 3162 IJ ლ — 1 759 1 - 3 m Contig |ORF | ID |ID | - 5 ~ _ ----ო 1067 | 2

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	Start Start (nt)	Stop	match acession	match gene name	mis &	* ident	length (nt)
1120	-	1 50	1 202	gi 142439	ATP-dependent nuclease (Bacillus subtilis)	64	30	153
1125		751	1 377	gi 581648	lepiB gene product (Staphylococcus epidermidis)	64	44	375
1688		1 402	1 214	pir A01365 TVMS	ltransforming protein K-ras - mouse	64 1	47	189
1 2472	-	2	1 358	gi 487282	Na+ -ATFase subunit J (Enterococcus hirae)	64	36	357
1 2989		1 520	1 356	gi 304134	largC (Bacillus stearothermophilus)	64 1	50	165
3013		1 630	1 352	gi 551699	oytochrome oxidase subunit I [Bacillus firmus]	64	51	279
1 3034		546	1 274	gi 1204349	hypothetical protein (GB:GB:D90212_3) [Haemophilus influenzae]	64	20	273
3197	-	613	308	gi 1009366		64 1	46	306
1 3303	-	06	1 362	gi 1107839	alginate lyase (Pseudomonas aeruginosa)	64	43	273
1 3852	2	82	1 288	gi 216746	D-lactate dehydrogenase [Lactobacillus plantarum]	64	42	207
1 3868	-		312	gi 149435	putative [Lactococcus lactis]	64	48	312
3918		099	331	gi 5532	acetyl-CoA acyltransferase [Yarrowia lipolytica]	64	46	330
1 4000	-	112	378	gi 984688	unknown [Saccharomyces cerevisiae]	64	44	267
4009		81	1 368	gi 39372	grsB gene product [Bacillus brevis]	64	41	288
4166		1 2	349	gi 149435	putative [Lactococcus lactis]	64	46	348
4366		5	1 307	gi 216267	ORF2 [Bacillus megaterium]	64	44	306
1 4457		- 2	1 400	gi 1197667	vitellogenin [Anolis pulchellus]	64	43	399
11	3	1539	2438	gi 438228	ORF C [Staphylococcus aureus]	63	32	1 006
24	7	5611	1 5423	gi 1369943	al gene product [Bacteriophage Bl]	63	34	189
53			390	gi 467441 	lexpressed at the end of exponential growyh under condtions in which he enzymes of the TCA cycle are repressed (Bacillus subtilis) gil467441 expressed at the end of exponential growyh under ondtions in which the enzymes of the TCA cycle are repressed Bacil	63	43	390
31	9	6329	5712	gi 496943	ORF [Saccharomyces cerevisiae]	63	47	618
1 44	123	114669	115019	pir A04446 QQEC	hypothetical protein F-92 - Escherichia coli	63	36	351
1 48	9	1 4403	6250	gi 43498	pyruvate synthase (Halobacterium halobium)	63	42	1848
50	2	1 3869	1 4738	gi 413967	lipa-43d gene product [Bacillus subtilis]	63	43	870
1 53	9	1 6764	1 5742	gi 474176	regulator protein (Staphylococcus xylosus)	63	49	1023

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig ORF	ORF ID	Start (nt)	Stop (nt)	match acession	match gene name	Eis &		l length (nt)
56	114	115880	117607	lgi1467409	DNA polymerase III subunit [Bacillus subtilis]	63	44	1728
57	┊Ξ.	7945	1376	gi 537036	IORF_0158 [Escherichia coli]	63	39	570
62	۳	2479	2114	gi 642656	Unknown [Rhizobium meliloti]	63	41	366
70	80	6562	1 7353	gi 1399821	PhoC [Rhizobium meliloti]	63	46	792
7.5	7	1 223	1 927	gi 149376	[HisG [Lactococcus lactis]	63	45	705
1.8	2	4912	4403	gi 413950	lipa-26d gene product [Bacillus subtilis]	1 63	42	510
91	2	904	1 7220	gi 466997	metH2; B2126_C1_157 [Mycobacterium leprae]	1 63	41	1857
1 91	8	10566	9448	gi 1204344	cystathionine gamma-synthase (Haemophilus influenzae)	63	45	1119
120		21	1508	gi 882657		63	46	1488
120	4	1 2722	1 4125	gi 665994	hypothetical protein [Bacillus subtilis]	63	34	1404
127	7	6064	1 7566	gi 40162	murE gene product (Bacillus subtilis)	63	44	1503
149	9	2321	1 2106	gi 148503	dnaK [Erysipelothrix rhusiopathiae]	63	40	216
149	126	110445	110170	gi 4870 	IORF 2, has similarity to DNA polymerase [Saccharomyces kluyveri] r[S15961[S15961 hypothetical protein 2 - yeast (Saccharomyces yveri) plasmid pSKL	63	42	276
164	7	507	1298	lgi1145476	CDP-diglyceride synthetase [Escherichia coli]	63	44	792
166	9	6066	8164	gi 151932	[fructose enzyme II [Rhodobacter capsulatus]	63	41	1746
169	4	1704	1886	gi 152886	elongation factor Ts (tsf) [Spiroplasma citri]	63	48	183
188	ر د	3145	2951	gi 1334547	GIY COI i14 grp IB protein [Podospora anserina]	63	42	195
195	113	111767	112804	gi 606100	ORF_0335 [Escherichia coli]	63	40	1038
201	2	607	1 2283	gi 433534 	arginyl-tRNA synthetase [Corynebacterium glutamicum] pir A49936 A49936 argininetRNA ligase (EC 6.1.1.19) - orynebacterium glutamicum	63	46	1677
206	114	15893	116489	gi 580828	N-acetyl-glutamate-gamma-semialdehyde dehydrogenase [Bacillus ubtills]	. 63	49	597
1 220	5	6977	1 5766	gi 216334	secA protein [Bacillus subtilis]	63	42	2004
1 221		74	1 907	gi 677945	AppA [Bacillus subtilis]	63	42	834
1 227	e -	944	1708	gi 1510558	cobyric acid synthase [Methanococcus jannaschii]	63	46	765
261	1 2	804	1070	gi 486511	ORF YKR054c [Saccharomyces cerevisiae]	63	45	267
269	2	3606	1960	gi 148221	DNA-dependent ATPase, DNA helicase [Escherichia coli] pirlJS0137 BVECRQ recQ protein - Escherichia coli	63	42	1647
	+							

Table

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	mis &	% ident	l length (nt)
278		7417	6176	gi 699273	CYSTATHIONINE GAMMA-SYNTHASE (Mycobacterium leprae) spiP468071METB_MYCLE CYSTATHIONINE GAMMA-SYNTHASE (EC 4.2.99.9) O-SUCCINYLHOMOSERINE (THIOL)- LYASE).	63	41	1242
1 287	- 5	738	1733	gi 405133		63	38	966
1 295		2	748	gi 1239983	hypothetical protein (Bacillus subtilis)	63	41	747
328	e 	1 2148	3134		Carrier protein (AA 1 - 437) Pseudomonas aeruginosa ir S11497 branched-chain amino acid transport protein braB - eudomonas aeruginosa	63	36	987
362	- 5	1626	1216	sp P35136 SERA_	D-3-PHOSPHOGLYCERAIE DEHYDROGENASE (EC 1.1.1.95) (PGDH).	63	38	411
404		326	1051	gi 1303816	YqeZ (Bacillus subtilis)	63	35	726
405	<u>س</u>	2101	1715	gi 1303914	YqhY (Bacillus subtilis)	63	42	387
1 406		451	722	gi 142152 	sulfate permease (gtg start codon) [Synechococcus PCC6301] pir A30301 GRYCS7 sulfate transport protein - Synechococcus sp. PCC 7942)	63	43	225
1 415	- 5	1 1048	2718	gi 1205402		63	41	1671
1 426	4	3575	2679	gi 393268	29-kiloDalton protein (Streptococcus pneumoniae) sp P42362 P29K_STRPN 29 KD MEMBRANE PROTEIN IN PSAA 5'REGION ORF1).	63	68 3	897
1 505	3	1347	2195	gi 1418999	orf4 [Lactobacillus sake]	63	40	849
1 507		1 2	574	gi 546917	comK [Bacillus subtilis, E26, Peptide, 192 aa]	63	35	573
1 562	- 5	146	1084	gi 43985	nifS-like gene (Lactobacillus delbrueckii)	63	45	939
675		1 427	1 215	gi 1510994	Serine aminotransferase [Methanococcus jannaschii]	63	29	213
989		3	1 230	gi 517356	Initrate reductase (NADH) [Lotus japonicus]	63	52	228
1 701		3	392	gi 881940	NorQ protein [Paracoccus denitrificans]	63	41	390
720	-	2	1 400	gi 47168	lopen reading frame [Streptomyces lividans]	63	35	399
179	-	571	1 287	gi 1261932	unknown [Mycobacterium tuberculosis]	63	41	285
1 907	- 1	1 22	321	gi 149445	ORF1 [Lactococcus lactis]	63	27	300
972		1 794	1 399	gi 1511235	[M. jannaschii predicted coding region MJ1232 [Methanococcus jannaschii]	63	27	396
1085		1154	618	gi 1204277	hypothetical protein (GB:U00019_14) [Haemophilus influenzae]	63	38	537
1094		3	1 542	gi 790943	urea amidolyase [Bacillus subtilis]	63	39	540
1108		e -	482	pir S49892 S498	regulation protein - Bacillus subtilis	63	44	480
1 1113		1231	617	gi 493017	endocarditis specific antigen [Enterococcus faecalis]	63	45	615

Table

204 243 249 195 291 210 216 174 693 192 225 402 132 189 396 399 393 360 168 195 312 303 990 | % ident | length | (nt) 46 30 34 46 47 33 20 35 36 40 25 21 40 47 48 37 48 54 57 30 36 44 21 8 sim 63 62 62 |carbamoyl-phosphate synthase (glutamine-hydrolysing) [Bacillus aldolyticus] HMG-CoA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir|A44756|A44756 | hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp. | Iproduct is similar to TnpA of transposon Tn554 from Staphylococcus ureus | [Clostridium butyricum] glutamate synthase (NADPH) [Azospirillum brasilense] pir|A49916|A49916 Pyruvate decarboxylase isozyme 2 (Swiss Prot. accession number P16467) [phosphoribosylformimino-praic ketoisomerase [Rhodobacter phaeroides] glutamate synthase (NADPH) (EC 1.4.1.13) - zospirillum brasilense |sp|P10537|AMYB |BETA-AMYLASE (EC 3.2.1.2) (1,4-ALPHA-D-GLUCAN MALTOHYDROLASE) |sp|P33940|YOJH_ |HYPOTHETICAL 54.3 KD PROTEIN IN ECO-ALKB INTERGENIC REGION |pir|A37967|A379 |neural cell adhesion molecule Ng-CAM precursor - chicken phenylacetaldehyde dehydrogenase [Escherichia coli] [flagellar MS-ring protein [Borrelia burgdorferi] (intrinsic membrane protein [Mycoplasma hominis] orf-1; novel antigen [Staphylococcus aureus] tagatose 6-P kinase [Streptococcus mutans] ORF360; putative [Bacteriophage LL-H] [Per6p gene product [Pichia pastoris] [pl00 protein [Borrelia burgdorferi] |rhoptry protein [Plasmodium yoelii] |pir|C33496|C334 |hisC homolog - Bacillus subtilis [amidase [Moraxella catarrhalis] |ATPase [Lactococcus lactis] | [Saccharomyces cerevisiae] |YrkA [Bacillus subtilis] |YqhY [Bacillus subtilis] [YbbF [Bacillus subtilis] (hMSH3 [Homo sapiens] | match gene name acession |gi|1303914 |gi|1184298 |gi|1041785 gi|1224069 |gi|1256135 gi (1321932 gi|1407784 gi|1480237 |qi|1303697 gi11256902 gi | 928989 gi|312443 gil1487982 gi|436132 |gi|836646 |gi|439126 |gi|623073 gi | 151259 |gi|296464 | 9 | 5985 | 6218 |gi|1490521 |gi|153675 1 2400 (nt) 695 1 250 204 245 1 227 402 1 293 135 1 252 1 217 338 399 395 1 176 195 1 308 193 1 195 360 1 400 171 312 361 | Start | | 3 | 1411 (nt) 2987 | 1 | 583 1 498 1 1 451 12 | 136 2994 | 1 | 266 1 1 | 440 609 1 1 793 1 1 720 338 ლ — 1 - 1 7 . — 9 ლ — 1 - 1 m Contig |ORF ID | ID -_ --- - _ _ _ _ 1300 2962 3043 3078 3139 3625 1325 1814 2021 2406 2961 3659 3783 2325 2335 3658 3900 4309 4367 4432 4468

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

1 1 2 10.02	Contig	IORE	Start (nt)	Stop (nt)	match acession	match gene name	E E I	% ident	l length (nt)
1856 1856 1860	37		1 2	1 721	gi 1107531	gene product [Campylobacter	62	33	720
12 1222 1730 141471234 OrdT [Haamophilas influences] 1 2 1222 1730 141471234 OrdT [Haamophilas influences] 1 2 1222 1730 141471234 OrdT [Haamophilas influences] 2 1222 1730 14175122 Highly Pydropholoic integral membrane procesn and phosphorylation Escherichis 62 23 1 1 2 1230 14175122 Highly Pydropholoic integral membrane procesn [Bacillus subtilis] 62 24 1 1 2 1220 14147053 14447053 14447054 14447056 14447054 14447056	38	115	110912	111589	gi 1222058		62	38	678
1 2 1539 1910 191173124 Ortil Hatemophilius inclinement of trivingent and phosphorylation Escherichia 62 35 15 15 15 15 15 15 15	:	125	119526	120329	gi 695280		62	41	804
1 2 555 Gill 1988174 Ellis Geneals of FPS-dependent Gat transport and phosphorylation Escherichia 62 35 188	57	- 5	1 2523	1780	gi 471234	orf1 [Haemophilus influenzae]	62	55	744
1 2 2 559 011 135132 httphily hydrophodoc integral membrane procean [Beaclines aubtills] 62 34 10 8250 9014 911470683 Shoop similarity with Arbbinding process 62 34 10 8250 9014 911470683 Shoop similarity with Arbbinding process 62 34 11 2 1309 91140083 101470683 101470683 101470683 101470683 101470683 101470684 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470683 101470684 101470683 101470683 101470683 101470683 101470683 101470684 101470683 101470684 10	57	6	6646	6350	gi 508174	EIIB domain of PTS-dependent Gat transport and phosphorylation Escherichia coli]	62	35	297
10 8220 9014 41470633 Shows similarity with ATP-binding proceins from other ABC-transport percens. 5 544 5 544 5 544 5 5	58		2	559		highly hydrophobic integral membrane protein (Bacillus subtilis) sp P42953 TAGG_BACSU TEICHOIC ACID TRANSLOCATION PERMEASE PROTEIN AGG.	62	34	558
1 1703 1704 [41140816 JactVM 4 gene product [Streeptomyces coelloolor] 62 44 44 47 47 47 47 47 4	67	100	8250	9014			62	34	765
1 1793 1320 gil139993 100P-N-acetylmuramoylalanine-D-glutamate ligase [Bacillus subtilis] 62 43 48 49 49 49 49 49 49 49	69	8	8315	7494	gi 46816	4	62	44	822
7 7034 9205 gil151007 phosphoglycerate dehydrogenase [Methanococcus Jannaachii] 62 42 42 1 2 520 gil151007 phosphoglycerate dehydrogenase [Methanococcus Jannaachii] 62 42 42 2 466 1068 gil153701 Alar-binding protein [Streptococcus memoniae 520 gil15365 mismatch repair protein [Streptococcus memoniae 520 37 4 6655 7562 gil1204866 L-fucose operon activator [Haemophilus influenzee] 62 37 5 6655 7562 gil1204866 L-fucose operon activator [Haemophilus influenzee] 62 37 6 6855 6004 gil85377 Appc [Bacillus subtilis] 62 37 8 6855 6004 gil85377 (product similar to E.coli PRP2 protein [Bacillus subtilis] 62 44 9 6855 6004 gil85377 (product similar to E.coli PRP2 protein [Bacillus subtilis] 62 44 1 24 554 gil467456 funknown [Bacillus subtilis] 62 33 1 24 554 gil467456 funknown [Bacillus subtilis] 62 41 1 24 554 gil46745 funknown [Bacillus subtilis] 62 41 1 24 554 gil46745 funknown [Bacillus subtilis] 62 62 62 1 24 662 6147 gil40336 phenylalanyl-RRA synthetase alpha subunit (Gly294 variant) unidentified 62 62 64 1 1 1225 614 gil40867 (Grobertical protein (SP:P20409 [Mycoplasma genitalium] 62 64 1 1225 614 gil104663 (Grobertical protein (SP:P20409 [Mycoplasma genitalium] 62 64 1 1225 614 gil104663 (Grobertical protein (SP:P202049) [Mycoplasma genitalium] 62 64 1 1225 614 gil40467 (Grobertical protein (SP:P202049) [Mycoplasma genitalium] 62 64 1 1225 614 gil40467 (Grobertical protein (SP:P202049) [Mycoplasma genitalium] 62 64 1 1225 614 gil40467 (Grobertical protein (GF:P202049) [Mycoplasma genitalium] 62 64 1 1225 614 gil40467 (Grobertical protein (GF:P202049) [Mycoplasma genitalium] 62 64 1 1 1225 614 614 614 614 614 614 614 614 614 614 614 614 614 614 614	80	e -	1 1793	1320		UDP-N-acetylmuramoylalanineD-glutamate ligase [Bacillus subtilis]	62	43	474
1 2 520 94 1531655 mismatch repair protein (Streptococcus pneumoniae) pir(28667)(228667 DNA 62 34 1 1 2 520 94 153655 mismatch repair protein hexA - Streptococcus neumoniae 62 37 34 1 1 2 520 94 153741 ATP-binding protein (Streptococcus mutans) 62 37 37 1 6855 5633 94 677947 Appc (Bacillus subtilis) 62 37 1 1 24 6823 5633 94 677947 Appc (Bacillus subtilis) 62 37 1 24 554 94 167756 1 1 24 554 94 167756 1 1 24 554 94 167756 1 1 24 554 94 167756 1 1 1 1 1 1 1 1 1	87		1 7034	9205	gi 217191		62	48	2172
1 2 520 gil153655 mismatch repair protein lexA - Streptococcus neumoniae 1 1 2 466 1068 gil153741 ATP-binding protein (Streptococcus mutans) 62 37 37 466 1068 gil153741 ATP-binding protein (Streptococcus mutans) 62 37 37 4 6825 7562 gil1204866 L-fucose operon activator [Haemophllus influenzae] 62 37 4 6823 5533 gil67747 Appc [Bacillus subtilis] 62 37 4 6823 5533 gil67747 Appc [Bacillus subtilis] 62 37 4 6823 6004 gil85377 Product similar to E.coli PRFA2 protein [Bacillus subtilis] 62 44 6823 6004 gil85377 Product similar to E.coli PRFA2 protein [Bacillus subtilis] 62 44 6823 6004 gil85377 Product similar to E.coli PRFA2 protein [Bacillus subtilis] 62 44 6823 614 615 614	100	۳ 	1 4051	3089	gi 1511047	phosphoglycerate dehydrogenase [Methanococcus jannaschii]	62	42	963
2 466 1068 91 153741 IATP-binding protein (Streptococcus mutans) 62 37 38 37 37 38 37 37 38 37 37	102		~	520	gi 153655	protein (Streptococcus pneumoniae)	62	34	519
7 6855 7562 91 1200866 IL-fucose operon activator [Haemophilus influenzee] 62 38 38 4 6823 5633 91 677947 Papec [Bacillus subtilis] 62 37 7 6825 6004 91 853777 Paroduct Similar to E.coil PRRZ protein [Bacillus subtilis] 62 44 7 7 7 7 7 7 7 7	112	7	1 466	1068	gi 153741	(ATP-binding protein (Streptococcus mutans)	62	37	603
4 6823 5633 gil677947 AppC [Bacillus subtilis] 62 37 8 6855 6004 gil85377 pir1855438 ywkE protein - Bacillus subtilis 62 44 9 6855 6004 gil85377 pir1855438 ywkE protein - Bacillus subtilis splP45873 HEMK_BACSU 62 44 1 24 554 gil467456 lunknown [Bacillus subtilis] 62 41 1 24 554 gil467456 lunknown [Bacillus subtilis] 62 41 1 24 554 gil467456 lunknown [Bacillus subtilis] 62 41 1 24 554 gil467456 lunknown [Bacillus subtilis] 62 41 1 24 554 gil467456 lunknown [Bacillus subtilis] 62 41 1 24 554 gil467456 lunknown [Bacillus sphaericus] 62 41 1 24 554 gil4067	114	7	1 6855	1 7562	gi 1204866	[L-fucose operon activator [Haemophilus influenzae]	62	38	708
8 6855 6004 gi 853777	116	4	6823	5633	gi 677947	AppC [Bacillus subtilis]	62	37	1191
1 24 554 gil467456 unknown [Bacillus subtilis]	124	& 	1 6855	1 6004	gi 853777 	protein (Bacillus su - Bacillus subtilis DASE (EC .3.3).	62	44	852
120 7591 6725 gi 1205807 Ireplicative DNA helicase [Haemophilus influenzae] 62 41 42 43 447	148		1 24	1 554	gi 467456	[Bacillus	62	50	531
3 1563 1153 9140067 X gene product [Bacillus sphaericus] 62 42 42 1156 14673 15632 91142219 P35 gene product (AA 1 - 314) [Escherichia coli] 62 38 62 38 62 1166 1447 911403936 Phenylalanyl-tRNA synthetase alpha subunit (Gly294 variant) unidentified 62 38 62 62 62 62 62 62 62 6		120	1 7591	1 6725		replicative DNA helicase [Haemophilus influenzae]	62	41	867
15 14673 15632 gi 42219 P35 gene product (AA 1 - 314) [Escherichia coli] 62 38	163	3	1 1503	1153	191140067		62	42	351
2 1166 1447 gi 403936 phenylalanyl-tRNA synthetase alpha subunit (Gly294 variant) unidentified 62 38	164	115	114673	115632	gi 42219	product (AA 1 -	62	38	096
2 2084 5089 gi 308861 GTG start codon [Lactococcus lactis] 62 44	165	5	1166	1447	gi 403936 	phenylalanyl-tRNA synthetase alpha subunit (Gly294 variant) unidentified cloning vector}	62	38	282
1 1225 614 gi 1046053 hypothetical protein (SP:P32049) [Mycoplasma genitalium] 62 41	166	7	1 2084	1 5089	gi 308861	[GTG start codon [Lactococcus lactis]	62	44	3006
	171		1 1225	614	gi 1046053	hypothetical protein (SP:P32049) [Mycoplasma genitalium]	62	41	612

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORE	Start (nt)	Stop (nt)	match acession	match gene name	e sim	8 ident	length (nt)
1 183		1 2521	1310	gi 143045	hemY [Bacillus subtilis]	62	45	1212
1 200		<u>ب</u>	956	gi 142439	ATP-dependent nuclease [Bacillus subtilis]	62	32	954
237	2	935	1966	gi 41695		62	44	1032
1 261	. –	4008	1 2605	gi 143121	ORF A; putative [Bacillus firmus]	62	1 42	1404
299	œ 	9477	4719	gi 467441 	lexpressed at the end of exponential growyh under conditions in which he enzymes of the TCA cycle are repressed [Bacillus subtilis] gi 467441 expressed at the end of exponential growyh under ondtions in which the enzymes of the TCA cycle are repressed Bacil	62	7.4	243
304	9	5018	3819	gi 153015	FemA protein [Staphylococcus aureus]	62	43	1200
324		2	262	gi 142717 	cytochrome aa3 controlling protein [Bacillus subtilis] pir[A33960[A33960] cta protein - Bacillus subtilis sp[P12946 CTAA_BACSU CYTOCHROME AA3 CONTROLLING PROTEIN.	62	30	261
325	7	1 269	1207	gi 581088	methionyl-tRNA formyltransferase [Escherichia coli]	62	39	939
1 332	9 -	1 4894	4631	lgi11499960	uridine 5'-monophosphate synthase [Methanococcus jannaschii]	62	36	264
1 355		5	370	gi 145925		62	32	369
365	8	1 6628	1 6804	gi 413943	lipa-19d gene product (Bacillus subtilis)	62	54	177
1 369	1 2	1 2744	1626	pir A43577 A435	1r A43577 A435 regulatory protein pfoR - Clostridium perfringens	62	42	1119
1 370		34	264	gi 40665		62	37	231
415	e -	1 2709	3176	gi 1205401	transport ATP-binding protein [Haemophilus influenzae]	62	35	468
429		1578	790	gi 1046024	Na+ ATPase subunit J [Mycoplasma genitalium]	62	40	789
444	7	1 704	1369	gi 581510 	nodulation gene; integral membrane protein; homology to Rhizobium eguminosarum nodI (Rhizobium loti)	62	37	999
477	7	751	1869	pir A48440 A484	ring-infected erythrocyte surface antigen 2, RESA-2 - Plasmodium falciparum	62	1 44	1119
1 485		1 241	1707	gi 17934	betaine aldehyd dehydrogenase [Beta vulgaris]	62	43	1467
1 487	<u>ه</u>	11141	1311	gi 149445	ORF1 [Lactococcus lactis]	62	31	171
1 494	1 2	1134	1313	gi 166835	ribulose bisphosphate carboxylase/oxygenase activase (Arabidopsis haliana)	62	37	180
518		193	882	gi 153491	O-methyltransferase (Streptomyces glaucescens)	62	1 39	069
534	2	1 369	2522	gi 1480429	putative transcriptional regulator [Bacillus stearothermophilus]	62	35	2154
551	9	1 4371	1 4820	gi 511113	ferric uptake regulation protein [Campylobacter jejuni]	62	37	450
1 574			570	gi 153000	enterotoxin B (Staphylococcus aureus)	62	43	570
								-

Table

828 435 477 345 246 819 675 366 171 159 165 399 213 411 246 372 594 591 201 231 231 255 243 297 183 | % sim | % ident | length (nt) 44 36 45 40 41 48 29 45 51 44 42 40 42 36 48 40 36 38 39 31 35 42 48 62 methyl coenzyme M reductase system, component A2 [Methanococcus jannaschii] | 10RF homologous to E.coli metB (Herpetosiphon aurantiacus) pir|S14030|S14030 | Hypothetical protein - Herpetosiphon aurantiacus fragment) acetyl coenzyme A acetyltransferase (thiolase) [Clostridium cetobutylicum] |HMG-COA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir|A44756|A44756 | hydroxymethyiglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp. M. genitalium predicted coding region MG181 [Mycoplasma genitalium] AgX-1 antigen [human, infertile patient, testis, Peptide, 505 aa] S. aureus - Putative coding regions of novel proteins similar to known proteins |homologous to N-acyl-L-amino acid amidohydrolase of Bacillus hypothetical protein (GB:X75627_4) [Haemophilus influenzae] |thioredoxin reductase [Eubacterium acidaminophilum] [alany1-tRNA synthetase [Methanococcus jannaschii] cell division inhibitor (Haemophilus influenzae) |deoxyuridine nucleotidohydrolase [Homo sapiens] |DNA-repair protein (recA) [Anabaena variabilis] epiB gene product [Staphylococcus epidermidis] [aspartokinase I [Methanococcus jannaschii] nontoxic component (Clostridium botulinum) ORF2, putative [Streptococcus pneumoniae] |nifS; B1496 C2 193 [Mycobacterium leprae] lipa-44d gene product [Bacillus subtilis] uroporphyrinogen III (Bacillus subtilis) |ipa-7d gene product [Bacillus subtilis] stearothermophilus (Bacillus subtilis) mrpC gene product [Proteus mirabilis] |Asparaginase [Bacillus licheniformis] |ORFC [Clostridium acetobutylicum] phnB protein [Escherichia coli] | match gene name acession gil1045865 |gi|1510649 |gi|1353197 |gi|1511613 |gi|1144332 gi | 1205451 gi | 1205822 |gi|1510641 |gi|1408501 |gi|475715 gi|147195 gi | 413931 |gi|581261 gi!485956 |gi|151259 |gi|688011 gi|460025 |gi|413968 1gi 1285708 gi | 710022 |gi|466883 |gi|142092 gi | 581648 1gi 140367 |gi|49272 Contig |ORF | Start | Stop ID | ID | (nt) | (nt) 1 1041 374 1491 414 830 1 478 348 944 400 1771 609 1 161 245 1 167 1 400 300 1 402 203 233 257 213 276 1 2 | 1715 344 1 493 1 331 823 31 1 1 396 1 1 | 692 1 267 1084 | 1 | 19 1 1 644 14 1898 1 1 35 1 1 463 ლ — 1 1 1 798 1 596 12 | 584 ٣ 3733 | 1 | 3 -1 - 3 - 1 - 5 -----_ _ -_ 2484 | 1 1071 1533 1537 2287 2386 3016 3116 3297 3609 3665 2490 1103 1217 590 655 656 919 687 864 700

Table?

| % ident | length (nt) # Sim |Similar to Saccharomyces cerevisiae SUA5 protein [Bacillus subtilis] | pir|S49358|S49358 ipc-29d protein - Bacillus subtilis sp|P39153|YWLC_BACSU | HYPOTHETICAL 37.0 KD PROTEIN IN SPOIIR-GLYC NTERGENIC REGION. !M. jannaschii predicted coding region MJ1083 [Methanococcus jannaschii] glutamine transport ATP-binding protein Q [Methanococcus jannaschii] mevalonate pyrophosphate decarboxylase [Saccharomyces cerevisiae] homologue to gene 30 (aa 1-59); putative [Bovine herpesvirus 4] hypothetical protein (GB:U00011_3) [Haemophilus influenzae] [hypothetical protein (SP:P33995) [Haemophilus influenzae] pir/C33356/C333 (prothymosin alpha homolog (clone 32) - human (fragment) amidophosphoribosyltransferase PurF [Rhizobium etli] [Na/H antiporter system ORF2 [Bacillus alcalophilus] |ORF95; putative [Lactococcus lactis phage BK5-T] exo-beta 1,3 glucanase [Cochliobolus carbonum] |ORFA gene product [Chloroflexus aurantiacus] ORF4; putative [Streptomyces violaceoruber] [hsdM protein (AA 1-520) [Escherichia coli] [hsdM protein (AA 1-520) [Escherichia coli] |tagatose 6-P kinase [Streptococcus mutans] |carbamate kinase [Haemophilus influenzae] lipa-34d gene product [Bacillus subtilis] |enterotoxin H [Staphylococcus aureus] |ORF2 [Clostridium perfringens] [YqeW [Bacillus subtilis] [ProX [Bacillus subtilis] |nikA [Escherichia coli] orf [Bacillus subtilis] |trac [Plasmid pAD1] match gene name acession |gi|1204846 |gi|1498756 |gi|1499931 |gi|1518679 | 4 | 1668 | 2531 |gi|1491643 gi11303813 1gi11510864 |gi|1205893 |gi|1205391 |gi|1109686 gi|1235684 |gi|1066504 |gi|153675 gi | 330705 gi1763513 |gi|928831 |gi|498839 1gi1388269 |gi|854656 |gi|466612 |gi|556881 gi|510692 |gi|41748 |gi|41748 1 5570 1 2243 1 303 1 223 Contig |ORF | Start | ID |ID | (nt) | 1 4288 1 4 | 3082 1 4605 1 2198 1 283 1 530 1 1 320 1 283 1 1 411 --1 1 1 11 12 11 | 1 -_ -... - æ — <u>б</u> - 2 _

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match acession	match gene name	as sim	% ident	length (nt)
132		1250	1 627	pir PQ0259 PQ02	hypothetical protein I - Enterococcus faecalis plasmid pAM-beta-1 (fragment)	61	43	624
1 149	6	3617	1 3075	gi 1144332	deoxyuridine nucleotidohydrolase [Homo sapiens]	61	40	543
149	122	1 8690	1 7869	gi 160047	p101/acidic basic repeat antigen [Plasmodium falciparum] pir(A29232 A29232 101K malaria antigen precursor - Plasmodium alciparum (strain Camp)	61	35	822
1 168		1 1915	2361		HIT protein, member of the HIT-family (Methanococcus jannaschii)	61	41	447
171	6 -	1 9675	1 7948	gi 467446	similar to SpoVB [Bacillus subtilis]	61	38	1728
1 174	- 3	1042	2340	gi 216374	glutaryl 7-ACA acylase precursor [Bacillus laterosporus]	61	49	1299
1 190	4	5034	4111	gi 409286	bmrU (Bacillus subtilis)	61	37	924
1 216		2	1 190	gi 415861	eukaryotic initiation factor 2 beta (eIF-2 beta) [Oryctolagus uniculus]	61	29	189
1 227	7	1 4161	5048	gi 216341	ORF for methionine amino peptidase (Bacillus subtilis)	61	41	888
1 238	4	1959	3047	gi 809543	CbrC protein {Erwinia chrysanthemi}	61	38	1089
1 247	-	7	1 694	gi 537231	ORF_f579 [Escherichia coli]	61	38	693
1 247	7	1 678	1 1034	gi 142226	chvD protein [Agrobacterium tumefaciens]	61	40	357
1 257	- 5	3523	1 2627	gi 699379	glvr-1 protein [Mycobacterium leprae]	61	40	1 897
1 268	- 5	3419	3051	gi 40364	ORFA1 [Clostridium acetobutylicum]	61	41	369
1 275	4	1 4621	1 4827	gi 1204848	hypothetical protein (GP:M87049_57) [Haemophilus influenzae]	61	36	1 207
772			1845	gi 784897	beta-N-acetylhexosaminidase [Streptococcus pneumoniae] pir[A56390[A56390] mannosyl-glycoprotein ndo-beta-N-acetylglucosaminidase (EC 3.2.1.96) precursor - treptococcus pneumoniae	61	45	1845
1 278	6	1 8003	1 7032	gi 467462		61	43	972
278	10	9878	8535	gi 1205919 	Na+ and Cl- dependent gamma-aminobutryic acid transporter [Haemophilus influenzae]	61	38	1344
1 283			1 366	lgi1755607	polyA polymerase [Bacillus subtilis]	61	36	366
1 288	- 5	1918	1496	gi 388108	cell wall enzyme [Enterococcus faecalis]	61	43	423
1 291		98	334	gi 454265	[FBP3 [Petunia hybrida]	61	38	249
318		1104	694	gi 290531 	similar to beta-glucoside transport protein [Escherichia coli] sp P31451 PTIB_ECOLI PTS_SYSTEM, ARBUTIN-LIKE IIB COMPONENT PHOSPHOTRANSFERASE_ENZYME II, B COMPONENT) (EC 2.7.1.69).	61	47	411
330	7	1912	1190	gi 1001805	hypothetical protein [Synechocystis sp.]	61	41	123

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORE	Start Start (nt)	Stop (nt)	match acession	match gene name	* sim	* ident	length (nt)
385	2	1513	1025	-+		61 1	42	489
426	-	794	1 399	gi 1303853	YqgF [Bacillus subtilis]	61	44	396
438	е.	810	1421	gi 1293660	AbsA2 [Streptomyces coelicolor]	61	36	612
454	-	1580	1 792	gi 733522	phosphatidylinositol-4,5-diphosphate 3-kinase [Dictyostelium iscoideum]	61	30	189
1 464	7	784	1 560	gi 1123120	C53B7.5 gene product [Caenorhabditis elegans]	61	38	225
470	8	1 6077	1 7357	gi 623073	ORF360; putative [Bacteriophage LL-H]	61	47	1281
605	-	554	1 279	gi 467484	unknown [Bacillus subtilis]	61	45	276
1 555	m — — -	1916	1 1296	gi 141800	anthranilate synthase glutamine amidotransferase (Acinetobacter alcoaceticus)	61	42	621
1 569	-	1711	1 857	gi 467090	B2235_C2_195 [Mycobacterium leprae]	61	47	855
1 585	7	961	1 803	sp P36686 SURE_	SURVIVAL PROTEIN SURE HOMOLOG (FRAGMENT).	61	33	159
1 592	<u> </u>	1694	1 1422	gi 1221602	immunity repressor protein [Haemophilus influenzae]	61	32	273
1 603	-	43	1 357	gi 507738	Hmp [Vibrio parahaemolyticus]	61	33	315
699		2467	1235	gi 1146243 	122.4% identity with Escherichia coli DNA-damage inducible protein; putative (Bacillus subtilis)	61	37	1233
675	e 	805	1101	gi 403373	[glycerophosphory] diester phosphodiesterase [Bacillus subtilis] pir[S37251[S3725] glycerophosphory] diester phosphodiesterase - acillus subtilis	61	36	297
1 703		1656	1 829	gi 537181	ORF_f470 [Escherichia coll]	61	32	828
1 728		1628	816	gi 806281	DNA polymerase I [Bacillus stearothermophilus]	61	39	813
821	-	61	318	1911709992	hypothetical protein [Bacillus subtilis]	61	38	258
826	1 5	2313	1 1567	1911609310	portal protein gp3 [Bacteriophage HK97]	61	40	147
923		1081	1 542	gi 143213		61	38	540
1124		59	370	gi 1107541	[C33D9.8 [Caenorhabditis elegans]	61	26	312
1492		548	1 276	gi 406397	Unknown [Mycoplasma genitalium	61	32	273
1602		46	318	gi 733522	[phosphatidylinositol-4,5-diphosphate 3-kinase [Dictyostelium iscoideum]	61	34	273
1 2500		775	1 290	gi 1045964	hypothetical protein (GB:U14003_297) [Mycoplasma genitalium]	61	31	288
2968	-	2	808	gi 397526	[clumping factor [Staphylococcus aureus]	61	55	807
3076		e -	248	gi 149373	[Lactococcus lactis]	61	41 -	246
	+		+					

171

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201

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270

1128 | % ident | length (nt) 39 42 20 41 46 47 45 24 48 42 46 20 38 20 32 47 20 43 61 61 61 61 61 61 61 61 61 61 09 % Sim 61 61 61 61 61 hsdM gene of Ecopril gene product [Escherichia coli] pir|s38437|s38437 hsdM | protein - Escherichia coli pir|s09629|s09629 hypothetical protein A - | Escherichia coli (SUB 40-520) beta-N-acetylhexosaminidase (Streptococcus pneumoniae) pir|A56390|A56390 |beta-N-acetylhexosaminidase [Streptococcus pneumoniae] pir|A56390|A56390 | mannosyl-glycoprotein ndo-beta-N-acetylglucosaminidase (EC 3.2.1.96) beta-N-acetylhexosaminidase [Streptococcus pneumoniae] pir/A56390/A56390 mannosyl-glycoprotein ndo-beta-N-acetylglucosaminidase (EC 3.2.1.96) beta-N-acetylhexosaminidase [Streptococcus pneumoniae] pir/A563901A56390 mannosyl-glycoprotein ndo-beta-N-acetylglucosaminidase (EC 3.2.1.96) beta-N-acetylhexosaminidase [Streptococcus pneumoniae] pir|A56390|A56390 beta-N-acetylhexosaminidase [Streptococcus pneumoniae] pir/A56390/A56390 large subunit of NADH-dependent glutamate synthase [Plectonema boryanum] mannosyl-glycoprotein ndo-beta-N-acetylglucosaminidase (EC 3.2.1.96) mannosyl-glycoprotein ndo-beta-N-acetylglucosaminidase (EC 3.2.1.96) mannosyl-glycoprotein ndo-beta-N-acetylglucosaminidase (EC 3.2.1.96) S. aureus - Putative coding regions of novel proteins similar to known proteins homologous to N-acyl-L-amino acid amidohydrolase of Bacillus |homologous to N-acyl-L-amino acid amidohydrolase of Bacillus | stearothermophilus [Bacillus subtilis] Respiratory nitrate reductase [Bacillus subtilis] ipa-29d gene product [Bacillus subtilis] stearothermophilus [Bacillus subtilis] precursor - treptococcus pneumoniae ribokinase [Escherichia coli] unknown (Bacillus subtilis) Yqew [Bacillus subtilis] AppD [Bacillus subtilis] |HrsA [Escherichia coli] HrsA [Escherichia coli] | match gene acession |gi|1303813 |gi|1408501 |gi|1009366 |gi|1339950 match |gi|1408501 |gi|413953 |gi|784897 gi|784897 |gi|677943 |gi|450688 |gi|147516 |gi|784897 1gi1976025 |gi|784897 |gi|784897 |gi|528991 1gi1976025 |gi|784897 1 5536 (nt) 1 220 383 329 345 336 1 233 1 273 172 1 227 401 1 740 442 401 274 402 268 398 Contig |ORF | Start | ID |ID | (nt) | 11477 1 6663 1 207 1 463 27 199 546 1 542 2 7 m m 7 - 2 - 1 -- _ -_ - _ 4038 | 1 9 3971 4374 4663 3609 3672 3724 4041 3662 3884 4047 4102 4155 4268 4389 4621 3728

738

441

219 396 399 381

303 273 402 345 336

195

| % sim | % ident | length 09 9 9 09 9 09 09 09 pir|A25526|A25526 ring-infected erythrocyte surface antigen recursor -Plasmodium falciparum (strain FC27/Papua New Guinea) sp|P13830|RESA_PLAFF RING-INFECTED ERYTHROCYTE SURFACE ANTIGEN RE (ring-infested erythrocyte surface antigen (Plasmodium falciparum) |carbonyl reductase (NADPH) [Rattus norvegicus] |para-nitrobenzyl esterase [Bacillus subtilis] [D-alanine racemase cds [Bacillus subtilis] |regulatory protein [Enterococcus faecalis] Ihypothetical protein [Synechocystis sp.] |ORF 4 [Saccharomyces kluyveri] |Orfl [Bacillus subtilis] | match gene name acession |gi|1001453 match |gi|1336656 |14 |11035 |10313 |gi|1217651 |gi|388109 | 6 | 3426 | 3725 | gi|410748 | 1 | 2047 | 1025 |gi|142822 | 4 | 2474 | 3607 | gi|468046 19114872 110814 | 9834 111917 | 112930 1 4 | 4364 | 4522 Contig |ORF | Start | Stop ID | ID | (nt) | (nt) 1 469 1 26 112 _ 113

111

16 33 37 39 1 41

723 1014

28

300

24

(nt)

S. aureus - Putative coding regions of novel proteins similar to known proteins

444 981 159

41 40 1023

47 39

43	4	1 2474	2474 3607 gi 46	gi 468046	para-nitrobenzyl esterase (Bacillus subtilis)	09	40	1134
44	110	6756	1 7769	gi 414234	thif [Escherichia coli]	1 09	52	1014
45	110	8874	1 9074	gi 343949	var1(40.0) [Saccharomyces cerevisiae]	1 09	44	201
56	118	127842	126430 gil46	gi 468764	mocR gene product [Rhizobium meliloti]	1 09	35	1413
09	- 5	173	388	gi 1303864	YqqQ (Bacillus subtilis]	09	33	216
63	7	357	1619	911467124	ureD; B229_C3_234 [Mycobacterium leprae]	09	43	1263
69		1 787	1 395	 gi 1518853	OafA (Salmonella typhimurium)	1 09	36	393
88			1188	gi 1480429	putative transcriptional regulator [Bacillus stearothermophilus]	1 09	30	1188
92	9	4735	1 3881	gi 349227	transmembrane protein [Escherichia coli]	1 09	37	855
92	7	5996	4923	gi 466613		09	38	1074
93		949	476	gi 1510925		09	27	474
96	9	7366	1 7578	gi 972715	accessory protein [Carnobacterium piscicola]	09	30	213
86	9	3212	1 4069	gi 467425	unknown [Bacillus subtilis]	09	42	828
102	10	7158	1 7430	7430 gi 143092 	acetolactate synthase small subunit [Bacillus subtilis] sp P37252 ILVN BACSU ACETOLACTATE SYNTHASE SMALL SUBUNIT (EC .1.3.18) (AHAS) (ACETOHYDROXY-ACID SYNTHASE SMALL SUBUNIT) (ALS).	09	37	273
109	111	9127	9127 10515 gi 12	gi 1255259	o-succinylbenzoic acid (OSB) CoA ligase [Staphylococcus aureus]	9	28	1389
109	112	110499	110499 11656	gi 141954		1 09 1	41	1158
119	5	1 4630	3134	gi 1524280	unknown [Mycobacterium tuberculosis]	09	45	1497
		1	1 1 1 1 1 1 1 1				+	+

| % sim | % ident | length 45 23 34 41 44 37 43 47 34 36 40 37 35 53 27 40 30 36 43 42 35 33 9 9 9 09 09 09 09 09 09 9 09 09 9 9 9 9 09 09 09 09 9 09 9 09 9 IORF3; PepY; putative oligoendopeptidase based on homology with Lactococcus |coded for by C. elegans cDNA ykl30e12.5; contains C2H2-type zinc fingers | [Caenorhabditis elegans] |similar to a B.subtilis gene (GB: BACHEMEHY_5) [Clostridium asteurianum] |hemagglutinin-neuraminidase fusion protein (Human parainfluenza irus 3) |sp|P37028|YADT_ |HYPOTHETICAL 29.4 KD PROTEIN IN HEML-PFS INTERGENIC REGION PRECURSOR. lactis PepF (GenBank Accession Number 232522) [Caldicellulosiruptor |NADP dependent leukotreine b4 12-hydroxydehydrogenase [Sus scrofa] S. aureus - Putative coding regions of novel proteins similar to known proteins |hypothetical protein (GB:U14003_130) [Haemophilus influenzae] hypothetical protein (GB:L19201_29) [Haemophilus influenzae] lhypothetical protein (GB:U00021_5) [Mycoplasma genitalium] glycine betaine transporter OpuD (Bacillus subtilis) [D9719.34p; CAI: 0.14 [Saccharomyces cerevisiae] (sp(P37347)YECD | HYPOTHETICAL 21.8 KD PROTEIN IN ASPS 5'REGION. 128.2 kDa protein [Streptococcus pneumoniae] |NaH-antiporter protein [Enterococcus hirae] |ipa-19d gene product [Bacillus subtilis] [clumping factor [Staphylococcus aureus] |ceuC gene product [Campylobacter coli] |murD gene product [Bacillus subtilis] |unknown [Mycobacterium tuberculosis] unknown [Mycobacterium tuberculosis] [putative [Bacillus subtilis] [HisD [Lactococcus lactis] |YqjX [Bacillus subtilis] |kdpA [Escherichia coli] | saccharolyticus] i match gene name |gi|1107529 |gi|1303975 |gi|1204532 |gi|1496003 |gi|1046160 |gi|1208889 |gi|1460077 |gi|1146230 |gi|1204366 |gi|1449288 |gi|1524397 |gi|1100737 |gi|148316 |gi|146547 |gi|580932 | 3 | 1331 | 2110 |gi|485280 |gi|413943 |gi|927798 |gi|332711 |gi|149377 1911431950 |gi|397526 2216 6013 1 6957 | 7646 11187 1 7866 1 2460 1 2415 112515 1410 1153 1409 1198 1 4769 1 4953 1 4573 1218 1 4595 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 1 703 4998 961 1 995 937 1 257 1 7704 114 | 8249 5919 2432 3804 5125 1 4082 1 4 | 5963 3641 13 13269 112234 1213 1 2809 1389 1 5696 159 65 11 | 511 1 860 ლ — 7 --6 7 - 5 _ - 7 _ 7 _ <u>б</u> 4 --9 ო <u>-</u> 140 145 158 159 173 198 201 203 206 214 237 241 260 264 267 291 150 157 160 161 212

1623 1011 993

933 855 282 198

780

1692

702 594 429 357 255 1029

384

1089

936 669 567 1215 1407

792

339

Table 7

S. aureus - Putative coding regions of novel proteins similar to known proteins

307 6 316 8 328 4						TARREST OF THE	11111111
-	1 3421	3176	 gi 1070014		1 09	36	246
-	1 4957	1 5823	gi 413952	lipa-28d gene product [Bacillus subtilis]	09	41	1 298
	1 2996	3484	gi 1204484		09	39	489
332 5	1 4887	1 4363	gi 1205449	colicin V production protein (pur regulon) [Haemophilus influenzae]	1 09	37	525
357 1	1 1062	1 532	gi 887842	single-stranded DNA-specific exonuclease [Escherichia coli]	09	41	531
375 2	96	362	gi 4857	adenylyl cyclase gene product [Saccharomyces kluyveri] r JQ1145 OYBYK adenylate cyclase (EC 4.6.1.1) - yeast ccharomyces kluyveri)	- 09	47	267
397 1	99	1 416	gil709999	Glucarate dehydratase [Bacillus subtilis]	09	37	351
409 1	1 2	163	gil499700	glycogen phosphorylase [Saccharomyces cerevisiae]	09	35	162
453 4	914	1237	gi 1196899	unknown protein [Staphylococcus aureus]	09	36	324
453 7	1 3838	1 3620	sp P12222 YCF1_	HYPOTHETICAL 226 KD PROTEIN (ORF 1901).	- 09	31	219
470 2	1 622	945	pir S30782 S307	pir S30782 S307 integrin homolog - yeast (Saccharomyces cerevisiae)	09	31	324
500 1	118	909	gi 467407	unknown [Bacillus subtilis]	- 09	36	489
503 3	1 752	1 982	gi 167835	myosin heavy chain [Dictyostelium discoideum]	09	34	231
505 4	1 2238	1 3563	gi 1510732	INADH oxidase [Methanococcus jannaschii]	09	26	1326
523 1	m 	1043	91/143331	(alkaline phosphatase regulatory protein [Bacillus subtilis] pir/A27650 A27650 regulatory protein phoR - Bacillus subtilis sp P23545 PHOR_BACSU ALKALINE PHOSPHATASE SYNTHESIS SENSOR PROTEIN HOR (EC 2.7.3).	09	41	1041
543 1		1 465	gi 1511103	cobalt transport ATP-binding protein O [Methanococcus jannaschii]	09	40	465
545 1		1 726	gi 1498192	putative [Pseudomonas aeruginosa]	- 09	40	726
556 1	2	1 1054	gi 1477402		09	42	1053
578 1	974	1 489	gi 1205129	H. influenzae predicted coding region HI0882 [Haemophilus influenzae]	- 09	42	486
594 1		624	gi 1212755	adenylyl cyclase {Aeromonas hydrophila}	09	45	624
604 1	m —	1 530	gi 145925		1 09	42	528
620 1	1 926	1 465	gi 1205483	bicyclomycin resistance protein [Haemophilus influenzae]	09	33	462
630 2	1 871	11122	gi 1486242	unknown [Bacillus subtilis]	- 09	41	252
645 2	1 574	1 425	gi 1205136	serine hydroxymethyltransferase (serine methylase) [Haemophilus influenzae]	09	28	150

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	:	Start	Stop	match	match gene name	+	+ % ident	length
a - +	a -	(nt)	(nt)	acession		- +	- +	(nt)
1 684		1 1082	843	gi 1205538	hypothetical protein (GB:U14003_302) [Haemophilus influenzae]	1 09	39	240
1 786		1 967	1 485	gi 1402944	orfRW1 gene product [Bacillus subtilis]	09	46	483
844		1 588	346	gi 790943	urea amidolyase [Bacillus subtilis]	1 09	40	243
851	.		1 726	gi 159661	GMP reductase [Ascaris lumbricoides]	1 09	41	726
871	1	1 1746	874	gi 1001493	hypothetical protein [Synechocystis sp.]	1 09	39	873
968		1558	839	gi 604926 	NADH dehydrogenase, subunit 5 [Schizophyllum commune] sp P50368 NU5M_SCHCO NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC .6.5.3).	09	39	720
806	- 5	1 448	1 753	gi 662880		1 09	31	306 1
979		2	1 595	gi 1429255	putative, orf1 [Bacillus subtilis]	- 09	30	594
1 1078		699	502	gi 581055 	inner membrane copper tolerance protein [Escherichia coli] gil811029 disulphide isomerase like protein [Escherichia coli] pir1847295 847295 inner membrane copper tolerance protein - scherichia coli	09	40	168
1112		1150	1 620	lgi1407885	ORF3 [Streptomyces griseus]	1 09	34	531
1135		484	1 275	gi 1171407	Vps8p [Saccharomyces cerevisiae]	1 09	36	210
1146		17	1 562	gi 1239981	hypothetical protein (Bacillus subtilis]	09	36	546
1291		1 716	1 360	pir S57530 S575	r S57530 S575 carboxyl esterase - Acinetobacter calcoaceticus	1 09	30	357
1 1332		1 336	169	gi 1222056	aminotransferase [Haemophilus influenzae]	1 09	44	168
1 1429			146	gi 1205619	[ferritin like protein [Haemophilus influenzae]	1 09	39	144
1722		570	1 286	gi 240052	dihydroflavonol-4-reductase, DFR [Hordeum vulgare=barley, cv. Gula, eptide, 354 aaj	09	36	285
2350	. – ;	385	1 200	gi 497626	ORF 1 [Plasmid pAQ1]	1 09	20	186
1 2936		1 519	310	gi 508981	prephenate dehydratase {Bacillus subtilis}	1 09	48	210
1 3027		1 568	302	gi 1146199		1 09	37	267
1 3084		1 20	1 208	gi 1407784	orf-1; novel antigen [Staphylococcus aureus]	1 09	51	189
1 3155		7	1 226	19111046097	cytadherence-accessory protein [Mycoplasma genitalium]	1 09	34	225
1 3603		368	186	gi 510108 	mitochondrial long-chain enoyl-CoA hydratase/3-hydroxycyl-CoA ehydrogenase alpha-subunit [Rattus norvegicus]	09	42	183
3665		1 486	244	gi 151259 	HMG-CoA reductase (EC 1.1.1.88) [Pseudomonas mevaloni]] pir[A44756 A44756 hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp.	1 09	42	243
3747		ж —	146	gi 474192	liucC gene product [Escherichia coli]	1 09	36	144
					. 5-1			

Table 7

| % sim | % ident | length (nt) |similar to trimethylamine DH [Mycoplasma capricolum] pir|S49950|S4950 | probable trimethylamine dehydrogenase (EC .5.99.7) - Mycoplasma capricolum (SGC3) (fragment) mitochondrial long-chain enoyl-CoA hydratase/3-hydroxycyl-CoA ehydrogenase |mitochondrial long-chain enoyl-CoA hydratase/3-hydroxycyl-CoA ehydrogenase | alpha-subunit [Rattus norvegicus] |peptide synthetase module [Microcystis aeruginosa] pir|S49111|S49111 | probable amino acid activating domain - icrocystis aeruginosa (fragment) unknown, similar to E.coli cardiolipin synthase (Bacillus subtilis)
! sp|P45860|YMIE_BACSU HYPOTHETICAL 58.2 PROTEIN IN NARI-ACDA NTERGENIC
! REGION. |chemoreceptor protein [Rhizobium leguminosarum bv. viciae] gi|780656 |pir|S48604|S486 |hypothetical protein - Mycoplasma capricolum (SGC3) (fragment) chemoreceptor protein [Rhizobium leguminosarum bv. iciae] |acyl-CoA-dehydrogenase [Streptomyces purpurascens] |novel antigen; orf-2 [Staphylococcus aureus] |Gat-1-P-DH, NAD dependent [Escherichia coli] transcription regulator (Bacillus subtilis) [PhoD precursor [Rhizobium meliloti] | alpha-subunit [Rattus norvegicus] [F59B2.3 [Caenorhabditis elegans] |VipB protein [Salmonella typhi] |BiP [Phaeodactylum tricornutum] |ORF_090a [Escherichia coli] pheB [Bacillus subtilis] ORF2 [Bacillus subtilis] [YqhT [Bacillus subtilis] |yeiH [Escherichia coli] [nikC [Escherichia coli] LORF F [unidentified] match gene name acession |gi|1488695 |gi|1399822 |gi|780656 |gi|1303901 |gi|405879 |gi|510108 |gi|426446 |gi|912461 1gi1602031 |gi|899317 gil508980 gi|510108 1gi1142833 |gi|912576 |gi|483940 |qi|508176 |gi|971345 |110 |14329 |15534 |gi|490328 |gi|536972 |gi|47382 1 2299 1 7460 Contig | ORF | Start | Stop ID | ID | (nt) | (nt) 1 6366 1 3523 1 495 1 280 1 2648 1 4651 1 4397 1 677 1 707 1 1 1 498 m _ -_ -е — _ = Ŋ –

Table ?

S. aureus - Putative coding regions of novel proteins similar to known proteins

96 1 1020 955 (41640201 Indication Indication Indication (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201 (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (41640201) (4164020000000000000000000000000000000000	Contig ORF	ORF	Start (nt)	Stop (nt)	match	match gene name	# sim	m % ident	length (nt)
1 2 820 911153822 Similar to B, subtilis baal (Bacillas subtilis) 59 82 82 82 82 82 82 82 8	68	1 2	1602	928		[Saccharomyces	5	9 1 32	1 645
1 2 2 820 Gill6722 ORF 5: putative [Bacilliae Subtili5] S9 1 1 1 1 1 1 1 1 1	96	4	1 4940	5473		protein of unknown function [Rhodobacter capsulatus]	5	9 1 33	1 534
1 166 1554 6115154 100F Dr. putetive (Bacillus filmus) 59 11 166 1554 6115154 100F Dr. putetive (Bacillus filmus) 59 11 12 12 13 13 14 12 13 14 14	86		1 2	820		B. subtilis DnaH (Bacillus	5	9 34	819
10 6214 6756 91115354 10187 55.9 Bacteriophage T4 518 11808 575 Bacteriophage T4 518 11808 575 Bacteriophage T4 518 518 5111086575 Bacteriophage T4 518 518 5111086575 Bact Reliance Campylobacter 19/4 11 518 518 511108657 Bact Indian subtilisis Campylobacter 19/4 518 518 51110855 Bact Indian subtilisis Bactilius abbtilisis Signatura Sig	119	- 1	166	1557		B; putative	, w	96 36	1392
16 12476 12351 9411006573 Bech [Rhizoblum mellioti] 59 1 1 1 1 1 1 1 1 1	1 120	110	6214	1 6756		ORF 55.9 [Bacteriophage T4]		66 39	543
1 306 195 191 1000	120	116	112476	113510		BetA [Rhizobium meliloti]		9 44	1035
1 370 645 9411256534 125.84 identity over 120 as with the Symenococcus 5pmraschii, purative 59 11001342 Pacifics Subcliss 18000hetical protein (SPFR2237) [Methanococcus 5pmraschii] 59 11001342 Phypothetical protein [Symechocystis sp.] 59 111001342 Phypothetical protein [Symechocystis sp.] 59 111001342 Phypothetical protein [Symechocystis sp.] 59 111001342 Phypothetical protein [Symechocystis sp.] 59 111001343 Phypothetical protein [Symechocystis sp.] 59 111001343 Phypothetical protein [Symechocystis sp.] 59 111001343 Phypothetical protein [Symechocystis sp.] 59 11101344 Phypothetical protein [Symechocystis sp.] 59 59 59 59 59 59 59 5	123	 	386	1 195		catalase [Campylobacter jejuni]	·	9 1 38	192
4 5278 5712 gill510635 hypothetical protein (SPP7057) [Methanococcus jannaschii] 59 1 1 3 509 gill01342 hypothetical protein [Synechocystis sp.] 59 1 1 1 1 1 1 1 1 1	1 130	 	370	1 645	gi 1256634	25.8% identity over 120 aa with the Synenococcus sp. MpeV protein; putative [Bacillus subtilis]	u)	9 31	276
1 3 509 [4111001342 [hypothetical protein [Symechocystis sp.]] 59 1 1 1 1 1 1 1 1 1	131	4 -	5278	5712	gi 1510655	hypothetical protein (SP:P42297) [Methanococcus jannaschii]		66 39	1 435
14 1529 2821 gill205165 hypothetical protein (SP:837764) [Haemophilus influencae] 59 1 19643 12176 gill1001381 hypothetical protein [Symechocyetis sp.] 59 1 1 1 1 1 1 1 1 1	164		. 3	1 509		hypothetical protein [Synechocystis sp.]	. ชา ! ! !	9 41	1 507
13 4127 3717 911104121 auxin-induced protein [Vigna radiate] 59 1 1 1 1 1 1 1 1 1	164	4	1529	2821		protein (SP:P37764)	<u>u</u> n	9 1 35	1293
1 12 12669 11503 941143036	164	119	1	121376				9 34	1734
1 2 1218 1688 911143036 Unidentified gene product [Bacillus subtilis] 59 1 1 12669 111503 911/62778 Niffs gene product [Anabaena azollae] 59 1 1 1 1 1 1 1 1 1	1 173	8	1 4727	1 3717		auxin-induced protein [Vigna radiata]	5	9 50	1011
12 12669 11503 1g1/762778 NifS gene product [Anabaena azollae] 59 1503 15103 1510240 1631519540 1631	179	1 2	1 2218	1688		[unidentified gene product [Bacillus subtilis]		9 33	531
5 4702 5670 gill510240 hemin permease [Methanococcus jannaschii] 59	195	112	112669	111503		NifS gene product [Anabaena azollae]	٤٦ -	9 41	1167
7 5719 6315 gil1511456	1 201	2	1 4702	1 5670		hemin permease [Methanococcus jannaschii]	٤)	9 32	696
1 102 461 gill204666 hypothetical protein (GB:X73124_53) [Haemophilus influenzae] 59 1 1 1 1 1 1 1 1 1	1 201	7 1	5719	6315		M. jannaschii predicted coding region MJ1437 [Methanococcus jannaschii]	<i>u</i> ,	9 1 34	1 597
3 1050 2234 gi 551531 12-nitropropane dioxygenase [Williopsis saturnus] 59 159 15 13293 4135 gi 1303709 YrkJ [Bacillus subtilis] 59 15 13078 13182 12167 gi 290489 Idfp (CG Site No. 18430) [Escherichia coli] 59 15 13078 13785 gi 149382 HisA [Lactococcus lactis] 15 13078 13182 1418551 Iransmembrane protein (kdpb) [Escherichia coli] 159 159 150 15	1 209	-	102	1 461	gi 1204666	hypothetical protein (GB:X73124_53) (Haemophilus influenzae)	٤,	9 42	1 360
5 3293 4135 9 11303709 YrkJ [Bacillus subtilis] 5 1 2167 9 1290489	214	3	1050	2234		2-nitropropane dioxygenase [Williopsis saturnus]	43 H	9 1 36	1185
2 3381 2167 gil290489 ldfp (CG Site No. 18430) [Escherichia coli] 59	214		3293	4135		[Bacillus	u)	9 1 32	1 843
5 3078 3785 gi 149382 HisA [Lactococcus lactis] 59	217	5 - 5	3381	2167			<u>u</u>)	9 44	1215
2 376 960 gi 1303791 YqeJ [Bacillus subtilis] 59	1 237	- 2	3078	3785		HisA [Lactococcus lactis]	<u>.</u>	9 1 38	1 708
1 1621 812 gil146551 transmembrane protein (kdpD) [Escherichia coli] 59	251	1 2	376	096		[Bacillus	<u>.</u>	9 34	1 585
5 4978 3860 gil405879 yeiH [Escherichia coli] 3 600 761 gil1303794 YqeM [Bacillus subtilis] 59	286		1621	812		[Escherichia	<u>.</u>	9 31	1 810
3 600 761 gi 1303794 YqeM [Bacillus subtilis]	316		1 4978	1 3860		yeiH [Escherichia coli]	<u>.</u>	9 1 32	1119
	370	۳ 	009	1 761		YqeM [Bacillus subtilis]	u) 	9 1 35	162

Table 2

| % sim | % ident | length (ut) |mannitol-phosphate dehydrogenase {Streptococcus mutans} pir|C44798|C44798 | mannitol-phosphate dehydrogenase MtlD - treptococcus mutans |bromoperoxidase BPO-A1 [Streptomyces aureofaciens] sp|P33912|BPA1 STRAU | NON-HAEM BROMOPEROXIDASE BPO-A1 (EC 1.11.1.-) BROMIDE PEROXIDASE) (BPO1) similar to eukaryotic Na+/H+ exchangers [Escherichia coli] | sp|P32703|XJCE_ECOLI HYPOTHETICAL 60.5 KD PROTEIN IN SOXR-ACS NTERGENIC | REGION (0549). IM. jannaschii predicted coding region MJ1577 [Methanococcus jannaschii] |iron(III) dicitrate transport ATP-binding protein FECE [Haemophilus (3R)-hydroxymyristol acyl carrier protein dehydrase [Haemophilus |bacterioferritin comigratory protein [Haemophilus influenzae] Igroup B oligopeptidase PepB (Streptococcus agalactiae [homoserine kinase homolog [Streptococcus pneumoniae] | 1354 | pir[A25620|A256 |staphylocoagulase - Staphylococcus aureus (fragment) |homologous to sp:HTRA_ECOLI [Bacillus subtilis] LukF-PV like component [Staphylococcus aureus] |pps1; B1496_C2_189 [Mycobacterium leprae] |hypothetical protein (Bacillus subtilis) |clumping factor [Staphylococcus aureus] ORF238 gene product [Porphyra purpurea] |ORF YAL022 [Saccharomyces cerevisiae] |trigger factor [Escherichia coli] |integrase [Bacteriophage phi-LC3] |orf3 [Haemophilus influenzae] |comE ORF3 [Bacillus subtilis] |ORF 3 [Spirochaeta aurantia] |unknown [Bacillus subtilis] match gene name influenzae (SUB 2-275) | influenzae acession |gi|1204610 |gi|1064809 |gi|1205311 |gi|1262748 |gi|1122758 gi | 1500472 |gi|1276668 |gi|1498295 |gi|1469939 |gi|1204511 |gi|147989 |gi|153746 1gi|396400 |gi|547513 |gi|152901 1gi | 709992 fgi|386681 gi | 293033 |gi|522150 |gi|289262 |gi|466882 1gi1397526 (nt) 1 723 Start | (nt) 1 2212 1 2836 1152 | 1 | 373 1 708 'n --Contig | ORF ID | ID - 2 -<u>ო</u> - 2 -ထ _ ~ _ _ ~ _

Table .

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	* sim	8 ident	length (nt)
1198		1 492	247	gi 142439	ATP-dependent nuclease [Bacillus subtilis]	59	1 26	246
1 1441		468	235	gi 1045942	glycyl-tRNA synthetase [Mycoplasma genitalium]	59	37	234
1 2103	1	1 1	186	gi 459250	triacylglycerol lipase [Galactomyces geotrichum]	59	33	186
1 2205		1 793	398	gi 1303794	YqeM [Bacillus subtilis]	59	38	396
1 2578	+ ·	484	284	qi 258003	insulin-like growth factor binding protein complex acid-labile ubunit [rats, liver, Peptide, 603 aa]	59	48	201
1 2967	2 -	145	348	gi 1212730	YqhK [Bacillus subtilis]	59	1 44	204
1 3012	1		1 248	gi 773571	neurofilament protein NF70 [Helix aspersa]	59	1 31	246
3544	1	۳ ا	4	gi 1055218	orotonase [Clostridium acetobutylicum]	59	42	399
3548		8	401	gi 1055218	crotonase [Clostridium acetobutylicum]	59	1 42	399
1 3580	1	869	1 351	gi 1055218	crotonase [Clostridium acetobutylicum]	59	42	348
1 3720		122	1 363	gi 1408494	homologous to penicillin acylase [Bacillus subtilis]	59	1 36	360
4171		<u>ب</u>	1 296	gi 1055218	crotonase [Clostridium acetobutylicum]	59	42	294
4305	-	618	310	gi 1524193	unknown [Mycobacterium tuberculosis]	59	39	309
 8 1		1242	622	gi 146913 	N-acetylglucosamine transport protein [Escherichia coli] pir B29895 WQEC2N phosphotransferase system enzyme II (Ec. 7.1.69), N-acetylglucosamine- specific - Escherichia coli sp P09323 PTAA_ECOLI PTS SYSTEM, N- ACETYLGLUCOSAMINE-SPECIFIC IIABC OMPONENT [EIIA	8 5	43	621
1 20	7 - 1	1 7020	5845	gi 50502	collagen alpha chain precursor (AA -27 to 1127) [Mus musculus]	58	20	1176
21	2	3234	3626	gi 1054860	phosphoribosyl anthranilate isomerase [Thermotoga maritima]	58	32	393
1 23	2	2841	1669	gi 1276880	EpsG [Streptococcus thermophilus]	58	29	1173
1 23	110	9301	0608	pir A31133 A311	r[A31133]A311 diaminopimelate decarboxylase (EC 4.1.1.20) - Pseudomonas aeruginosa	58	37	1212
38	129	122555	122884	lgi1973249	vestitone reductase [Medicago sativa]	58	37	330
44	-	~	1 406	gi 289272	ferrichrome-binding protein [Bacillus subtilis]	58	33	405
45			552	gi 29464	embryonic myosin heavy chain (1085 AA) [Homo sapiens] ir S12460 S12460 myosin beta heavy chain - human	58	33	552
1 55	2 -	759	1 538	gi 158852	glucose regulated protein [Echinococcus multilocularis]	58	32	222
1 62	113	8493	8908	gi 975353	Kinase-associated protein B [Bacillus subtilis]	58	35	426
63	e 	1553	1717	gi 166926	[Arabidopsis thaliana unidentified mRNA sequence, complete cds.], ene product [Arabidopsis thaliana]	58	32	165
+								

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	# sim	% ident	length (nt)
1 67	113	112017	111229	gi 1228083	INADH dehydrogenase subunit 2 [Chorthippus parallelus]	28	41	1 687
96	80	8208	9167	gi 709992	hypothetical protein (Bacillus subtilis)	28	42	096
107	2	1 2065	1364	gi 806327	[Escherichia coli hrpA gene for A protein similar to yeast PRP16 and RP22 [Escherichia coli]	28	37	702
112		4519	5613	gi 155588 	glucose-fructose oxidoreductase [Zymomonas mobilis] pir A42289 A42289 glucose-fructose oxidoreductase (EC 1.1) recursor - Zymomonas mobilis	288	38	1095
114	9	7318	1 6503	gi 1377843	unknown [Bacillus subtilis]	85	38	816
143	7	1 2261	1395	pir A45605 A456	mature-parasite-infected erythrocyte surface antigen MESA - Plasmodium falciparum	5.88	31	867
151	- 5	717	1 950	 gi 1370261	unknown [Mycobacterium tuberculosis]	28	31	234
154	9	6015	1 4627	gi 1209277	IPCTHom1 gene product [Chlamydia trachomatis]	28	41	1389
154	116	114281	113541	gi 146613	DNA ligase (EC 6.5.1.2) [Escherichia coli]	28	39	741
155	۳ 	1 2269	1892	gi 1303917	YqiB (Bacillus subtilis)	28	34	378
174		1056	1 529	gi 904198	hypothetical protein [Bacillus subtilis]	28	26	528
189	4	1533	1769	gi 467383	DNA binding protein (probable) [Bacillus subtilis]	28	25	237
201	ю —	1 2669	1 3307	gi 1511453	endonuclease III [Methanococcus jannaschii]	28	34	639
1 208		7	1 238	gi 1276729	phycobilisome linker polypeptide [Porphyra purpurea]	28	1 29	237
1 220	11.	114575	13058	gi 397526	clumping factor [Staphylococcus aureus]	58	1 51	1518
1 231	<u>م</u>	1629	1 1474	gi 1002520	MutS [Bacillus subtilis]	58	1 45	156
1 233	9	4201	1 3497	gi 1463023	No definition line found [Caenorhabditis elegans]	28	39	705
1 243	110	9303	110082	gi 537207	ORF_f277 [Escherichia coli]	58	32	780
1 257		331	11143	gi 1340128	ORF1 (Staphylococcus aureus)	58	44	813
302	1 2	1 460	801	gi 40174	ORF X [Bacillus subtilis]	28	34	342
1 307	111	1 6984	6127	gi 1303842	YqfU (Bacillus subtilis]	58	30	828
321	۳ -	1914	1 2747	gi 1239996	hypothetical protein [Bacillus subtilis]	58	41	834
1 342	4	1 2724	3497	gi 454838	ORF 6; putative [Pseudomonas aeruginosa]	28	41	774
348			663	gi 467478	unknown [Bacillus subtilis]	58	36	663
401	5	1 384	1 605	gi 143407	[para-aminobenzoic acid synthase, component I (pab) [Bacillus ubtilis]	58	53	222

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig ID	IORF	Start (nt)	Stop (nt)	match acession	match gene name	& sim	% ident	length (nt)
437		1 325	1554	gi 1303866	YqgS [Bacillus subtilis]	28	35	1230
445		105	1442	gi 581583	protein A [Staphylococcus aureus]	288	32	1338
453	. –	189	1 965	gi 1009455	unknown [Schizosaccharomyces pombe]	58	34	177
453	- 5	2748	1 2047	gi 537214	lyjjG gene product [Escherichia coli]	58	40	702
479	7	731	1444	gi 1256621	126.7% of identity in 165 aa to a Thermophilic bacterium hypothetical protein 6; putative [Bacillus subtilis]	58	36	714
490		606	547	gi 580920	InddD (gtaA) polypeptide (AA 1-673) [Bacillus subtilis] pir/S06048 S06048 probable rodD protein - Bacillus subtilis sp Pl3484 TAGE BACSU PROBABLE POLY(GLYCEROL-PHOSPHATE) LPHA-GLUCOSYLTRANSFERASE (EC 2.4.1.52) (TECHOIC ACID BIOSYNTHESIS ROTEIN E).	ω 	39	363
517	-		1164	sp1P47264 Y018_	HYPOTHETICAL HELICASE MG018.	28	30	1164
517	9	4182	4544	lgi1453422	orf268 gene product [Mycoplasma hominis]	58	29	363
546	m 	1 2802	4019	gi 886052	restriction modification system S subunit [Spiroplasma citri] gi 886052 restriction modification system S subunit [Spiroplasma itri]	28	37	1218
562	-	e -	179	gi 43831	InifS protein (AA 1-400) [Klebsiella pneumoniae]	288	34	177
009	- 5	1347	1156	gi 1183839	unknown [Pseudomonas aeruginosa]	288	48	192
604	- 5	1231	1001	gi 1001353	hypothetical protein [Synechocystis sp.]	28	41	231
619		1	504	gi 903748	lintegral membrane protein (Homo sapiens)	58	43	504
625		2	364	gi 1208474	hypothetical protein [Synechocystis sp.]	58	43	363
635		1492	1 755	gi 1510995		28	41 1	738
645			846	gi 677882 	ileal sodium-dependent bile acid transporter [Rattus norvegicus] gi 677882 ileal sodium-dependent bile acid transporter [Rattus orvegicus]	58	33	846
645	m —	906	1556	gi 1239999	hypothetical protein [Bacillus subtilis]	28	41	651
665		177	1 532	gi 1204262	hypothetical protein (GB:L10328 61) [Haemophilus influenzae]	28	39	240
674	1	635	1 327	gi 498817	ORF8; homologous to small subunit of phage terminases [Bacillus ubtilis]	58	39	309
675	- 5	1312	908	gi 42181	osmC gene product [Escherichia coli]	288	28	507
745	- 1	618	310	gi 1205432	coenzyme PQQ synthesis protein III (pqqIII) [Haemophilus influenzae]	58	32	309
799	7	242	1174	gi 1204669	collagenase [Haemophilus influentae]	28	36	933
800	7	1096	614	gi 171963 	TRNA isopentenyl transferase (Saccharomyces cerevisiae) sp P07884 MOD5_YEAST TRNA ISOPENTENYLTRANSFERASE (EC 2.5.1.8) ISOPENTENYL- DIPHOSPHATE: TRNA ISOPENTENYLTRANSFERASE) (IPP RANSFERASE) (IPPT).	58	37	483
	1				+11111111111111111111111111111111111111	+		

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

******	1	A-manual and	A-community		T			+
Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	% sim	% ident	length (nt)
854		1108	1 605	gi 466778	lysine specific permease [Escherichia coli]	288	44	504
885	-	1 481	1 242	gi 861199	Incotoporphyrin IX Mg-chelatase subunit precursor [Hordeum vulgare]	28	33	240
891		е -	1 527	gi 1293660	AbsA2 [Streptomyces coelicolor]	58	31	525
942		931	1 467	gi 405567	traH [Plasmid pSK4]	28	30	465
1002		1 952	521	gi 577649		28	34	432
1438			261	gi 581558 	isoleucyl frwa synthetase (Staphylococcus aureus) sp P41368 SYIP STAAU ISOLEUCYL-TRNA SYNTHETASE, MUPIROCIN RESISTANT EC 6.1.1.5) (ISOLEUCINE TRNA LIGASE) (ILERS) (MUPIROCIN RESISTANCE ROTEIN).	23	30	261
1442	·	2	1 463	gi 971394	similar to Acc.No. D26185 [Escherichia coli]	288	34	462
1873	-	1 480	241	gi 1339951	small subunit of NADH-dependent glutamate synthase [Plectonema boryanum]	28	38	240
1876		m 	158	gi 529216	No definition line found [Caenorhabditis elegans] sp P46503 YLX7_CAEEL HYPOTHETICAL 7.3 KD PROTEIN F23F12.7 IN HROMOSOME III.	58	33	156
1989	-	108	401	gi 1405458	YneR [Bacillus subtilis]	28	29	294
2109		е -	1 401	gi 1001801	hypothetical protein [Synechocystis sp.]	58	31	399
2473		1 288	1 145	gi 510140	[ligoendopeptidase F [Lactococcus lactis]	58	38	144
2523		1 452	1 228	gi 644873	catabolic dehydroquinate dehydratase [Acinetobacter calcoaceticus]	58	37	225
3041		2	211	gi 1205367	oligopeptide transport ATP-binding protein (Haemophilus influenzae)	58	39	210
3094	-		1 263	gi 1185288	isochorismate synthase [Bacillus subtilis]	58	38	261
3706		m 	1 383	gi 456614	mevalonate kinase [Arabidopsis thaliana]	288	48	381
3854			1 402	gi 808869		58	32	402
4082		51	1 224	gi 508551 	ribulose-1,5 bisphosphate carboxylase large subunit -methyltransferase [Pisum sativum]	28	37	174
4278		е 	206	gi 180189 	cerebellar-degeneration-related antigen (CDR34) [Homo sapiens] gi 182737 cerebellar degeneration-associated protein [Homo sapiens] pir[A29770 A29770 cerebellar degeneration-related protein - human	28	37	204
19	7	1 7818	1 7363	gi 1001516	hypothetical protein [Synechocystis sp.]	57	31	456
23	111	1 9663	1 8872	gi 606066	ORF £256 [Escherichia coli]	57	29	792
31		1 4801	1 2402	gi 153146		57	32	2400
38	114	111611	110796	gi 144859	ORF B [Clostridium perfringens]	57	31	816
46	114	112063	13046	gi 1001319	hypothetical protein [Synechocystis sp.]	57	25	984
		!!!!!!!			+ 4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

-						-		•
Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	* sim	% ident	length (nt)
51	3	1411	1187	pir B33856 B338	hypothetical 80K protein - Bacillus sphaericus	57	38	225
54			453	gi 684950	staphylococcal accessory regulator A [Staphylococcus aureus]	57	31	453
1 75		3	1 239	gi 1000470	C27B7.7 [Caenorhabditis elegans]	57	42	237
92	- 5	1 3855	3061	gi 143607	sporulation protein (Bacillus subtilis)	57	35	795
96	E	1 4006	4773	gi 144297	acetyl esterase (XynC) [Caldocellum saccharolyticum] pir1B372021B37202 acetylesterase (EC 3.1.1.6) (XynC) - Caldocellum accharolyticum	57	34	768
107	- 3	1480	1 2076	gi 460955	TagE [Vibrio cholerae]	57	42	597
109		1 5340	1 5933	gi 1438846	Unknown [Bacillus subtilis]	57 1	41	594
112	6 -	6299	1 7701	gi 1486250	Unknown [Bacillus subtilis]	57 1	33	1023
114	4	6384	4108	gi 871456	putative alpha subunit of formate dehydrogenease [Methanobacterium hermoautotrophicum]	57	37	2277
126	- 5	430	1053	gi 288301	ORF2 gene product [Bacillus megaterium]	57	37	624
131	- 2	1 6537	729	9111511160	IM. jannaschii predicted coding region MJ1163 [Methanococcus jannaschii]	57	38	261
133	Э	1 2668	1 2201	91/1303912	YqhW [Bacillus subtilis]	57	40	468
133	4	1 3383	1 2784	gi 1221884	(urea?) amidolyase (Haemophilus influenzae)	57	37	009
147	4	1 2164	1694	gi 467469	unknown [Bacillus subtilis]	57	33	471
160	1 2	1 1293	1 1060	gi 558604	chitin synthase 2 [Neurospora crassa]	57	28	234
163	8	1 5687	4764	gi 145580	rarD gene product [Escherichia coli]	57	38	924
168	9 -	4336	5325	gi 39782	133kDa lipoprotein (Bacillus subtilis)	57	32	066
170	2	1 3297	3455	gi 603404	Yer164p [Saccharomyces cerevisiae]	57	37	159
1 221	9	1 8026	6089	gi 1136221	carboxypeptidase [Sulfolobus solfataricus]	57	32	1218
228	m 	1348	1791	gi 288969	fibronecin binding protein [Streptococcus dysgalactiae] pir S33850 S33850 fibronecin-binding protein - Streptococcus ysgalactiae	57	32	444
1 263	4	4411	3686	gi 1185002	dihydrodipicolinate reductase [Pseudomonas syringae pv. tabaci]	57	42	726
1 276		494	255	gi 396380	No definition line found [Escherichia coli]	57	40	240
1 283	- 5	1 335	1324	gi 773349	BirA protein [Bacillus subtilis]	57	32	066
1 297	-	1 469	1 236	gi 1334820	reading frame V [Cauliflower mosaic virus]	57	46	234
342	m —	1993	1 2805	gi 1204431	hypothetical protein (SP:P33644) [Haemophilus influenzae]	57	35	813

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

	1111						+	+
Contig	IORE	Start (nt)	Stop (nt)	match acession	match gene name	ae Eis	% ident	length (
375	9	3340	3741	gi 385177		57	26	402
1 433	9	3286	4011	gi 1524117	alpha-acetolactate decarboxylase [Lactococcus lactis]	57	40	726
470	. —	1 903	1145	gi 804819	protein serine/threonine kinase [Toxoplasma gondii]	57	30	243
487	5	1391	1723	gi 507323	ORF1 [Bacillus stearothermophilus]	57	28	333
498	-	1 274	852		NADH-ubiquinone oxidoreductase subunit 4L [Podospora anserina]	57	34	579
1 503		343	173	gi 1502283		57	30	171
1 505	7	1619	1284	gi 466884	B1496_C2_194 [Mycobacterium leprae]	57	40	336
519	7	1182	1 2549	gi 1303707	YrkH (Bacillus subtilis)	57	34	1368
522	7	3234	1945	gi 1064809	homologous to sp:HTRA_ECOLI [Bacillus subtilis]	1 57	36	1290
538	7	606	1415	gi 153179 	phosphorinothyrcin n-acetyltransferase [Streptomyces coelicolor] pir JH0246 JH0246 phosphinothricin N-acetyltransferase (EC 2.3.1) Streptomyces coelicolor	57	40	507
547		896 1	1 486	gi 467340		57	1 05	483
599		1062	532	sp P20692 TYRA_	PRREPHENATE DEHYDROGENASE (EC 1.3.1.12) (PDH).	57	41	531
1 620	- 5	757	572	gi 1107894	unknown [Schizosaccharomyces pombe]	57	38	186
622	- 5	1600	1130	gi 173028	thioredoxin II [Saccharomyces cerevisiae]	57	39	471
625	7	362	1114	gi 1262366	hypothetical protein [Mycobacterium leprae]	57	34 1	753
089		→	204	gi 143544 	RNA polymerase sigma-30 factor [Bacillus subtilis] pir1A28625 A28625 transcription initiation factor sigma H - acillus subtilis	57	30	204
069		m	629	gi 466520	pocR [Salmonella typhimurium]	57	29	627
969	-	1 2	433	gi 413972	lipa-48r gene product (Bacillus subtilis]	57	33	432
704		36	638	gi 1499931	[M. jannaschii predicted coding region MJ1083 [Methanococcus jannaschii]	57	36	603
732		1 2316	1621	gi 1418999		57	37	969
1 746		451	1 227	gi 392973	[Rab3 [Aplysia californica]	57	42	225
1 757		20	466	gi 43979 	L.curvatus small cryptic plasmid gene for rep protein [Lactobacillus rvatus]	57	45	447
1 862	. . .	2	295	lgi11303827	[YqfI [Bacillus subtilis]	57	21	294
1049		1 907	455	gi 1510108		57	35	453
1117		1387	1 695	gi 896286	NH2 terminus uncertain [Leishmania tarentolae]	57	28	693
	11111							

Table 2

| % ident | length (nt) H & SIM |histidine utilization repressor G (Klebsiella aerogenes) pir/A36730|A36730 | hutG protein - Klebsiella pneumoniae (fragment) sp/P19452/HUTG KLEAE | FORMIMINOGLUTAMASE (EC 3.5.3.8) FORMIMINOGLUTAMATE HYDROLASE) (HISTIDINE | UTILIZATION PROTEIN G) FRAGMENT). glucose-fructose oxidoreductase (EC 1.1. -. -) recursor - Zymomonas mobilis |HMG-CoA reductase (EC 1.1.1.88) [Pseudomonas mevalonii] pir|A44756|A44756 | hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88) Pseudomonas sp. tetracenomycin C resistance and export protein (Streptomyces laucescens) IM. jannaschii predicted coding region MJ1143 [Methanococcus jannaschii] |restriction modification enzyme, subunit M1 [Methanococcus jannaschi1] glucose-fructose oxidoreductase [Zymomonas mobilis] pir|A42289|A42289 glutamate synthase (ferredoxin) [Synechocystis sp.] pir|S46957|S46957 |voltage-activated calcium channel alpha-1 subunit (Rattus orvegicus) |leukotoxin secretion ATP-binding protein [Haemophilus influenzae] (EC 1.4.7.1) - ynechocystis sp. pir|JV0037|RDEC |nitrate reductase (EC 1.7.99.4) alpha chain - Escherichia coli |Pristinamycin I synthase I [Streptomyces pristinaespiralis] putative membrane protein; putative [Bacillus subtilis] lisochorismate synthase [Haemophilus influenzae] |T06C10.5 gene product (Caenorhabditis elegans) |ipa-29d gene product [Bacillus subtilis] |1-phosphofructokinase [Escherichia coli] |uracil permease [Escherichia coli] glutamate synthase (ferredoxin) |SpoIIIAG [Bacillus subtilis] |ORF 0464 [Escherichia coli] [patatin [Solanum tuberosum] [YqgF [Bacillus subtilis] |ORF4 [Bacillus subtilis] |EF [Streptococcus suis] |yeiH [Escherichia coli] match gene name acession |gi|1303853 |gi|1511146 gi|1237015 gi|1000365 |gi|1511222 gi|1204540 |gi|1086633 |gi|1205301 gi11483603 gil310083 gi | 142780 19i1313777 |gi|882472 gi|153490 gil155588 |gi|413953 |gi|151259 |gi|298032 gi|405894 gi | 515938 |gi|149204 gi1405879 gi|21512 (nt) 1 242 1 256 (nt) 1 746 1 468 1 793 ო ლ — Contig |ORF ID |ID - 5 _ -_ _ _ --_

Table 2

s. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF 11D	Start (nt)	Stop (nt)	match acession	match gene name	e sim	8 ident	length (nt)
22	· - -	1 4248	1 5177	gi 132222		56	33	1 086
38	128	121179	122264	gi 1480705	lipoate-protein ligase [Mycoplasma capricolum]	56	34	1086
44	<u>۾</u>	1 1861	1 2421	gi 490320	Y gene product [unidentified]	26	1 31	561
44	115	110103	110606	gi 1205099	hypothetical protein (GB:L19201_1) [Haemophilus influenzae]	95	39	504
20	9	1 4820	5161	gi 209931	fiber protein [Human adenovirus type 5]	26	48	342
53	4	1 2076	1 2972	gi 623476	transcriptional activator [Providencia stuartii] sp P43463 AARP_PROST TRANSCRIPTIONAL ACTIVATOR AARP.	26	30	897
	9	1 5656	6594	gi 466613	nikB [Escherichia coli]	1 56	32	1 686
6.8	m 	1 2364	1810	gi 482922		1 56	39	555
96		1 203	913	gi 145594	cAMP receptor protein (crp) [Escherichia coli]	1 56	35	711
109	121	118250	117846	gi 1204367	hypothetical protein (GB:U14003_278) [Haemophilus influenzae]	26	1 27	405
112	œ 	5611	1 6678	gi 155588	Iglucose-fructose oxidoreductase [Zymomonas mobilis] pir A42289 A42289 glucose-fructose oxidoreductase (EC 1.1) recursor - Zymomonas mobilis	26	40	1068
131	۳ 	1 6404	1 5100	gi 619724	MgtE [Bacillus firmus]	95	1 30	1305
138		1 65	1 232	gi 413948	ipa-24d gene product (Bacillus subtilis)	96	1 31	168
138	4	823	1521	gi 580868	lipa-22r gene product [Bacillus subtilis]	95	31	669
146	7	1 740	447	gi 1046009	M. genitalium predicted coding region MG309 [Mycoplasma genitalium]	26	1 37	294
149	7	1639	1067	gi 945380	terminase small subunit [Bacteriophage LL-H]	1 56	35	573
163		7	223	gi 143947	glutamine synthetase [Bacteroides fragilis]	56	30	222
166	- 5	6745	1 6449	gi 405792	ORF154 [Pseudomonas putida]	95	1 26	297
187		1 31	1 393	gi 311237	[H(+)-transporting ATP synthase [Zea mays]	56	30	363
190		7	1 373	gi 1109686	ProX [Bacillus subtilis]	26	35	372
191	8	111538	1 9943	gi 581070	acyl coenzyme A synthetase [Escherichia coli]	56	35	1596
195		1291	1 647	gi 1510242	collagenase [Methanococcus jannaschii]	56	34	645
230	۳ 	1 2323	1 2072	gi 40363	heat shock protein [Clostridium acetobutylicum]	1 56	39	252
238	ر د	1 3383	1 3775	gi 1477533	sarA {Staphylococcus aureus}	1 56	31	393
270	7 -	813	1712	gi 765073		26	41	006
	i +-					1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

290 1 1221 1620 1612 161	ig	ORF	Start (nt)	Stop (nt)	match acession	match gene name		Eis 	* ident	length (nt)
1 12.94 12.79 641 1211155 W. Jannacachi I predicted coding region WNISC Patronacachi Jannacachi 1 12.94 12	1 290		1 3221	1632	ı —		 	56	34	1590
1 1779 [4] [1001801] hypothetical protein [Synechcoystis sp.] 56 31 2 1779 [6] [145356] [101] Gene product [Rhitothum millotil] 56 26 26 2 1762 [8] [145304] [1-ribblioklame [Borderichha coll]] 56 39 1 4 1772 200 [9] [11010054] [Pypothetical protein [Synachcoystis applied]] 56 39 31 4 1772 200 [9] [11010054] [Pypothetical product [Borderich Molycoylase [Borderich	1 297	- 5	1140	1373	gi 1511556	M. jannaschii predicted coding region MJ1561 [Methanococcus jannaschii]	<u> </u>	26 1	40	234
1 1752 1279 1271 1270 127	321	- 5	1 2947	1799		hypothetical protein [Synechocystis sp.]	-	26	31	1149
1 176 182 191101330 Il-ribulokiaase [Escherichia coli] 186 182 191101330 Il-ribulokiaase [Escherichia coli] 186 182 191101330 Il-ribulokiaase [Escherichia spila] 182 192 191101330 Il-ribulokiaase [Escherichia spila] 182 191101330 Il-ribulokiaase [Escherichia spila] 182 191101330 Il-ribulokiaase [Escherichia subtilis] 182 191101330 Il-ribulokiaase [Escherichia subtilis] 182 1911101330 Il-ribulokia influence 182 182 191101330 Il-ribulokia influence 182 192	359	- 5	1279	641		noll gene product [Rhizobium meliloti]		26	26	639
1 1920 1229 54110319904 Phytochetical procein (Synechocystis sp.) 56 51 51 51 52 52 52 52 52	371	- 5	1 360	1 1823		L-ribulokinase [Escherichia coli]	<u> </u>	56	39	1464
1 1900 192 gillid36994 5-Hrdi receptor [Homo sapiens] 56 48 48 48 48 48 48 48 4	391	4	1762	1 2409		hypothetical protein [Synechocystis sp.]		26	34	648
1 1907 1203 q:1114008466 H874A gene product [Bacillus subtilis] 56 31 32 32 32 32 32 32 32	1 402		1 380	192		5-HT4L receptor [Homo sapiens]		56 1	48	189
1 1307 1017 1911203194	416	4	1 2480	1 2109		HS74A gene product [Bacillus subtilis]		56	31	372
1 1907 1017 6411205194	1 424	8 -	1756	2334		acetolactate decarboxylase [Bacillus subtilis]		56	32	579
2 2423 1812 191147661 Lap78 Fasteurella haemolytica] 56 37 181 182 2152 1233 1911170661 Lap78 Fasteurella haemolytica] 56 37 181 1	457		1 1907	1 1017		formamidopyrimidine-DNA glycosylase [Haemophilus influenzae]		56	36	891
1 1 1284 911217049 DrnO protein [Salmonella typhimurium] 56 37 1 1 1 1284 911217049 DrnO protein [Salmonella typhimurium] 56 37 1 1 1 1 1 1 1 1 1	458	1 2	1 2423	1 1812		terminase [Bacteriophage SPP1]		26 1	37	612
1 1 1284 gil217049 brnQ procein [Salmonella typhimurium] 56 37 1 1 1 1284 gil467109 irin; 308 Ribosomal protein SiB alamine acetyltransferase; 229_Cl_170 56 43 1 1 1 1 1 1 1 1 1	1 504	- 5	1 2152	1283		Lpp38 [Pasteurella haemolytica]		26	38	870
3 1099 1701 191467109 Irim, 308 Ribosomal protein SIB alanine acetyltransferase; 229_C1_170 56 43 1 1 1 1 1 1 1 1 1	511	-		1284		brnQ protein [Salmonella typhimurium]		26 -	37	1284
5 3547 3774 gji1229106 IXK930.1 (Caenorhabditis elegans)	604	<u>۾</u>	1099	1701	_	rim; 30S Ribosomal protein S18 alanine acetyltransferase; 229_C1_170 [Mycobacterium leprae]		26	43	603
1 35 400 gil153929 NADPH-sulfice reducatase flavoprotein component [Salmonella yphimurium] 56 38 38 38 38 38 38 38 3	099	2 -	1 3547	3774		[ZK930.1 [Caenorhabditis elegans]		56	30	228
2 1385 1095 91 1510801 Inydrogenase accessory protein [Methanococcus jannaschii] 56 35 35	707	-	35	1 400		NADPH-sulfite reducatase flavoprotein component [Salmonella yphimurium]		56 -	38	366
1 1 495 gil413948 Ippa-24d gene product [Bacillus subtilis] 56 35 35	1 709	- 1	1 1385	1095		hydrogenase accessory protein [Methanococcus jannaschii]		56 -	38	291
1 87 677 gi 928836	718	-	1	495	_	lipa-24d gene product [Bacillus subtilis]		56	35	495
1 776 399 gil1511513 ABC transporter, probable AFP-binding subunit [Methanococcus jannaschii] 56 33	744	-	1 87	1 677		repressor protein [Lactococcus lactis phage BK5-T]		56 1	35	591
1 3 407 gi 12205382 cell division protein (Haemophilus influenzae) 56 34 19 930 gi 1222161 permease (Haemophilus influenzae) 56 28	190		1 776	399		transporter, probable ATP-binding subunit	_	56 -	33	378
1 19 930 gil1222161 permease [Haemophillus influenzae] 56 28	795		e -	1 407		cell division protein [Haemophilus influenzae]		26	34	405
1 3 515 gi 1256621 126.7% of identity in 165 aa to a Thermophilic bacterium hypothetical 56 33 31 1 2 466 gi 547513 lorf3 [Haemophilus influenzae] . 56 37 2 1049 732 gi 886022 MexR [Pseudomonas aeruginosa] 56 31	813		19	1 930		permease (Haemophilus influenzae)		56	28	912
1 2 466 gil547513 lorf3 [Haemophilus influenzae] . 56 37	855		e 	515		26.7% of identity in 165 aa to a Thermophilic bacterium hypothetical protein 6; putative [Bacillus subtilis]		56	33	513
2 1049 732 gi 886022 MexR [Pseudomonas aeruginosa]	896		5	1 466		orf3 [Haemophilus influenzae]	-	56	37	465
1 5 223 gi 184251 HMG-1 [Homo sapiens]	973	- 1	1049	1 732		MexR [Pseudomonas aeruginosa]		26 –	31	318
	1 1203		S	1 223		[HMG-1 [Homo sapiens]		26 -	34	219

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	e sin	8 ident	length (nt)
1976		1 452	237	gi 9806 	Iysine-rich aspartic acid-rich protein [Plasmodium chabaudi] r S22183 S22183 lysine/aspartic acid-rich protein - Plasmodium baudi	95 -	33	216
2161		2	1 400	gi 1237015	ORF4 (Bacillus subtilis)	1 56	27	399
2958		362	1 183	gi 466685	No definition line found [Escherichia coli]	1 56	26	180
2979		421	1 212	gi 1204354	Spore germination and vegetative growth protein [Haemophilus influenzae]	1 56	40	210
2994	7	1 526	326	gi 836646	phosphoribosylformimino-praic ketoisomerase [Rhodobacter phaeroides]	1 56	29	201
3026		179	1 328	gi 143306	penicillin V amidase [Bacillus sphaericus]	1 56	30	150
3189	-	1 289	146	gi 1166604	Similar to aldehyde dehydrogenase [Caenorhabditis elegans]	1 56	37	144
3770		1 63	1 401	gi 1129145	acetyl-CoA C-acyltransferase [Mangifera indica]	1 56	43	339
4054	7	1 720	361	gi 1205355	Na+/H+ antiporter (Haemophilus influenzae)	1 56	31	360
4145			1 324	gil726095	long-chain acyl-CoA dehydrogenase [Mus musculus]	1 56	36	324
4200		1 505	254	gi 155588	glucose-fructose oxidoreductase [Zymomonas mobilis] pir A42289 A42289 glucose-fructose oxidoreductase (EC 1.1) recursor - Zymomonas mobilis	95 -	40	252
4273	-	675	355	gi 308861	GTG start codon [Lactococcus lactis]	1 56	33	321
-4	т — —	4095	3436	gi 5341 	Putative orf YCLX8c, len:192 [Saccharomyces cerevisiae] r S53591 S53591 hypothetical protein - yeast (Saccharomyces evisiae)	55	25	099
11	112	1 9377	8505	gi 21673	haloacetate dehalogenase H-1 [Moraxella sp.]	52	32	873
12	4	5133	1 4534	gi 467337	unknown (Bacillus subtilis]	55	26	900
19	2	5404	1 5844	gi 1001719	hypothetical protein (Synechocystis sp.)	1 55	25	441
23	13	114087	112339	gi 474190	iucA gene product [Escherichia coli]	55	30	1749
32		5368	1 6888	19111340096	unknown [Mycobacterium tuberculosis]	1 55	37	1521
34	m 	1 2569	1808	gi 1303968	YqjQ [Bacillus subtilis]	55	39	762
34	5	1 3960	3412	gi 1303962	YqjK (Bacillus subtilis)	55	33	549
36		1291	647	gi 606045	ORF_0118 [Escherichia coli]	55	27	645
36	9	6220	5243	gi 1001341	hypothetical protein [Synechocystis sp.]	55	31	978
47	<u>ش</u>	1 3054	1 3821	gi 1001819	hypothetical protein (Synechocystis sp.]	1 55	21	768
49		2065	1127	gi 403373 	glycerophosphoryl diester phosphodiesterase [Bacillus subtilis] pir[S37251 S37251 glycerophosphoryl diester phosphodiesterase - acillus subtilis		36	939
	+	+						

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig ORF	ORF	Start (nt)	Stop (nt)	match acession	match gene name	& sim	\$ ident	length (nt)
69	111	9968	1 9565	gi 153053		55	23	009
75	8	881	1273	 gi 41698	L-histidinol: NAD+ oxidoreductase (EC 1.1.1.23) (aa 1-434) scherichia coli]	55	33	393
82	6	115387	114194	gi 1136221	carboxypeptidase [Sulfolobus solfataricus]	55	35	1194
87	4	3517	4917	gi 1064812		55	26	1401
88	1 2	1172	1636	gi 882463		55	35	465
95	- 1	127	516	gi 1377832		55	36	390
100	- 5	836	1 2035	gi 1370274		55	36	1200
100	- 5	1 5137	1 4658	gi 396660		55	29	480
108	- 3	1 4266	1 2986	gi 1499866	M. jannaschii predicted coding region MJ1024 [Methanococcus jannaschii]	55	31	1281
114	3	1 2616	1 1834	gi 1511367		55	29	783
144	- 3	1 1805	1476	gil1100787	unkown [Saccharomyces cerevisiae]	55	35	330
165	- 5	6212	5508	gi 1045884	M. genitalium predicted coding region MG199 [Mycoplasma genitalium]	55	27	705
1 189	- 5	1 2205	1 2576	gi 142569	ATP synthase a subunit [Bacillus firmus]	55	35	372
191	9 -	9136	1 6857	gi 559411		55	39	1 2280
194	- 5	1 364	636	gi 1145768		55	34	273
500	4	1335	1676	gil473357		55	35	342
211	2 -	1693	1145	gi 410130	ORFX6 [Bacillus subtilis]	55	37	549
213	2 -	644	1372	gi 633692		55	28	729
214	7	4144	5481	gi 1001793	Inypothetical protein [Synechocystis sp.]	55	30	1338
221	7	111473	9197	gi 466520	pocR [Salmonella typhimurium]	55	32	722
233	8 - -	1 5908	1 4817	gi 1237063	unknown [Mycobacterium tuberculosis]	55	38	1092
1 236	1 4	1 1375	1 2340	gi 1146199	putative [Bacillus subtilis]	55	32	996
1 243	1 2	1 380	1885	lgi1459907		55	1 29	1506
1 258	- 1	1 786	394	gi 455006		55	36	393
281		126	938	gi 1408493		55	35	813
316		1323	2102	gi 1486447	[LuxA homologue [Rhizobium sp.]	55	30	780
1 326	- 5	1 2968	2744	gi 1296824	proline iminopeptidase [Lactobacillus helveticus]	55	36	225
	+-		+					

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
351	7	2322	1 1429	gi 1204820	hydrogen peroxide-inducible activator [Haemophilus influenzae]	55	1 28	894
353	4	2197	2412	gi 1272475	chitin synthase [Emericella nidulans]	55	1 50	216
380	-	14	379	gi 142554	ATP synthase i subunit [Bacillus megaterium]	55	37	366
383		462	1 232	gi 289272	ferrichrome-binding protein (Bacillus subtilis)	55	36	231
386			938	gi 1510251	DNA helicase, putative [Methanococcus jannaschii]	55	30	1 986
410	2	1208	1 1891	gi 1205144		55	1 27	684
483	2	411	833	gi 413934	ipa-10r gene product [Bacillus subtilis]	55	1 26	423
529	3	7771	1433	gi 606150	ORF_f309 [Escherichia coli]	55	33	345
555	7	1088	585	gi 143407	para-aminobenzoic acid synthase, component I (pab) [Bacillus ubtilis]	55	1 28	504
565	- 1	402	1 202	gi 1223961	CDP-tyvelose epimerase [Yersinia pseudotuberculosis]	1 55	41	201
582		751	1 452	gi 1256643	20.2% identity with NADH dehydrogenase of the Leishmania major mitochondrion; putative [Bacillus subtilis]	52	36	300
645	<u>-</u>	1 2260	2057	gi 210824 	fusion protein F (Bovine respiratory syncytial virus) pirjJ01481 VGNZBA fusion glycoprotein precursor - bovine espiratory syncytial virus (strain A51908)	. 55	52	204
672	2	1 957	2216		M. jannaschii predicted coding region MJ1322 [Methanococcus jannaschii]	55	36	1260
730	-	955	479	gi 537007	IORF_f379 [Escherichia coli]	55	30	477
737	-	1859	945	gi 536963	CG Site No. 18166 [Escherichia coli]	1 55	30	915
742	7	1 228	1 572	gi 304160	product unknown [Bacillus subtilis]	55	38	345
817	7	1211	1 903	gi 1136289	histidine kinase A [Dictyostelium discoideum]	1 55	29	309
819	-	1 582	355	gi 558073	polymorphic antigen [Plasmodium falciparum]	55	22	228
832	- 2	1152	724	gi 40367	ORFC [Clostridium acetobutylicum]	55	32	429
840		1 769	1 386	gi 1205875	pseudouridylate synthase I [Haemophilus influenzae]	55	39	384
1021		23	1 529	gi 48563	 beta-lactamase [Yersinia enterocolitica]	55	38	507
1026		09	1 335	gi 47804	Opp C (AA1-301) [Salmonella typhimurium]	55	1 26	276
1525			1 282	gi 1477533		55	29	282
1814	- 1	1 224	985	gi 1046078		55	38	762
3254		427	1 254	gi 413968	lipa-44d gene product [Bacillus subtilis]	55	30	174
		!!!!!!	1	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,				

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

	-							+
Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	e sim	% ident	length (nt)
3692		989	345	gi 216773		1 55	32	342
3721	1		312	gi 42029	ORF1 gene product [Escherichia coli]	55	31	312
3799		m —	1 272	gi 42029	ORF1 gene product [Escherichia coli]	55	38	270
3889		1 22	1 423	gi 1129145		55	45	402
3916		2	385	gi 529754	spec [Streptococcus pyogenes]	55	38	384
3945		4	1 198	gi 476252	phase 1 flagellin (Salmonella enterica)	55	36	195
4074	7	1 488	1 246	gi 42029	ORF1 gene product [Escherichia coli	55	38	243
4184		7	1 343	gi 1524267		52	28	342
4284		14	1 208	gi 1100774	ferredoxin-dependent glutamate synthase [Synechocystis sp.]	55	36	195
4457	2	644	378	gi 180189 	cerebellar-degeneration-related antigen (CDR34) [Homo sapiens] gi[182737 cerebellar degeneration-associated protein [Homo sapiens] pir[A29770[A29770] cerebellar degeneration-related protein - human	55	88	267
4514	-	2	1 244	gi 216773		55	32	243
4599	-	432	1 217	gi 1129145	acetyl-CoA C-acyltransferase [Mangifera indica]	55	42	216
4606		416	1 210	gi 386120	(myosin alpha heavy chain (S2 subfragment) (rabbits, masseter, eptide Partial, 234 aa]	55	27	207
5	80	5348	4932	1911536069	ORF YBL047c [Saccharomyces cerevisiae]	54	27	417
12	7	1 7166	6165	gi 1205504	homoserine acetyltransferase [Haemophilus influenzae]	54	30	1002
	116	117086	115326	gi 474192	liucC gene product [Escherichia coli]	54	31	1761
35		2	979	gi 48054 	small subunit of soluble hydrogenase (AA 1-384) [Synechococcus sp.] ir S06919 HQYCSS soluble hydrogenase (EC 1.12) small chain - nechococcus sp. (PCC 6716)	54	36	978
37	111	9437	1 8667	1911537207	ORF_f277 [Escherichia coli]	54	38	771
37	112	8165	8332	gi 1160967	palmitoyl-protein thioesterase [Homo sapiens]	54	37	168
46	115	13025	113804	gi 438473	protein is hydrophobic, with homology to E. coli ProW; putative Bacillus subtilis]	54	28	780
56	7	1 203	1 736	gi 1256139	YbbJ [Bacillus subtilis]	54	34	534
57	113	111117	110179	gi 1151248	linosine-uridine preferring nucleoside hydrolase (Crithidia fasciculata)	54	32	939
99	1 2	516	1133	gi 1335781	[Cap [Drosophila melanogaster]	54	29	618
70	110	8116	8646	gi 1399823	PhoE [Rhizobium meliloti]	54	31	531
	i !							-

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

15 12556 11801 18p1P029831TCR S IETRACYCLINE RESISTANCE PROTEIN. 1 1 1005 1289 91 1064811	match match gene name acession		a sin	% ident	length (nt)
5 4915 5706 9111064811 function u 1 3005 12289 9111205366 01190eptt 1 2 13585 12095 911143727 putative 1 4 2337 2732 91 143727 putative 1 4 2337 2732 91 143727 putative 1 2 1720 2493 91 144297 acetylest 1 2 1720 1336 91 42473 pyruvate 1 2 1525 1172 91 1377834 unknown [Electric	P02983 TCR_S		54	29	756 1
4 3005 2289 91 1205366 01190pept 12 2596 1556 91 710495 protein ki 12 3585 2095 91 143727 putative 14 2337 2732 91 143727 putative 15 1600 2493 91 144297 acetyl est 12 1720 2493 91 144297 acetyl est 12 1555 1172 91 1377834 unknown 18 19 4831 5469 91 903305 ORF73 Bac 13 1369 13 137882 phosphate 16 3263 4543 91 1240916 prolyl-tR 16 1363 4543 91 1240916 prolyl-tR 16 1363 4550 91 1240916 prolyl-tR 16 5683 4250 91 1240916 prolyl-tR 18 19208 19720 91 1240916 R09EIO.3 18 19208 19720 91 1240916 R09EIO.3 19 19 19 19 19 19 19 1	1064811 function unknown	-	54	33	192
2 3586 1556 91710495		ilus influenzae]	54	33	1171
2 3585 2095 91143727	1710495		54	33	1041
4 2337 2732 91153724 MALC [Street 1 2 1720 2493 91114297 acetyl est 1 2 1720 2493 91144297 acetyl est 1 2 525 1172 91 42473 pyruvate 1 2 525 1172 91 1377834 unknown [6 694 7251 91 1204976 phosphate 6 3263 4543 91 1204976 prolyl-tR 1 1 1 1 1 1 1 1 1	143727 putative [Bacillus		54	30	1491
2 1720 2493 gi 144297 acetyl est 5 1600 3306 gi 42473 pyruvate of 2 525 1172 gi 1377834 unknown [IIIII G694 7251 gi 1511039 phosphate 6 3263 4543 gi 1504976 prolyl-tRR 7 12062 22243 gi 143582 spoiliEA 8 18208 19720 gi 143582 spoiliEA 9 1909 1905 gi 1467378 unknown [IIIII IIIII IIIII IIIIII IIIIIII	MalC [Streptococcus		54	41	396
5 1600 3306 9142473 Pyruvate of the control of the contr	144297 acetyl esterase (XynC) Caldocellum acetylesterase (EC 3.1.1.6) (XynC)	saccharolyticum] pir B37202 B37202 - Caldocellum accharolyticum	54	34	174
2 525 1172 91 1377834 Unknown	192473 pyruvate oxidase [Escherichia		54	36	1707
9 4831 5469 91903305 ORF73 [Bac 13263 4543 91 1204976 Prolyh-tRR 120 22243 91 1204976 Prolyh-tRR 120 22243 91 143582 SpoIIIEA 120 120208 19720 91 436965 [malA] ger 12 1090 1905 91 1240016 R09E10.3 12 1090 1905 91 1240016 R09E10.3 13 12655 13059 pir(S00485 S004 9ene 11-1 13 12655 13059 pir(S00485 S004 9ene 11-1 13 12655 14219 91 62964 arylamine 17 4141 4701 91 682769 mccE gene 10 6994 8742 91 41951 ip-27d ger 13 13377 2214 91 490328 LORF F [un	1377834 unknown [Bacillus		54	23	648
13 6694 7251 911511039 phosphate 6 3263 4543 911204976 proly1-tRR 120 121602 122243 911143582 spoilifeA 120 121602 122243 911143582 spoilifeA 120 19720 911240016 R09E10.3 122 1090 1905 911467378 unknown [E 1 1322 663 911353761 myosin II 1312 663 911353761 myosin II 1312 663 911353761 myosin II 132 663 911303813 Yqqw Bacc 17 5055 4219 9162964 arylamine 17 4141 4701 9162964 mccE gene 10 6994 8742 91413951 ipa-274 gene 11 1337 12214 91490328 LORF F [uncapped 10 10 10 10 10 10 10 1	1903305 ORF73 [Bacillus		54	28	639
6 3263 4543 gi 1204976	1511039 phosphate	anococcus jannaschii]	54	32	558
120	11204976	_	54	34	1281
6 5683 4250 gi 436965 18 19208 19720 gi 1240016 2 1090 1905 gi 467378 1 1322 663 gi 1353761 3 2030 3709 gi 1303813 7 5055 4219 gi 62964 7 5055 4219 gi 62964 7 4141 4701 gi 882769 10 6994 8742 gi 43951 3 3377 2214 gi 490328			54	32	642
18 19208 19720 gi 1240016 1809E10.3 (Caenorhabditis elega 2 1090 1905 gi 467378 Unknown (Bacillus subtilis) 1 1322 663 gi 1353761 myosin II heavy chain (Naegler 13 12655 13059 pir 500485 5004 gene 11-1 protein precursor 3 2030 3709 gi 1303813 YqeW (Bacillus subtilis) 7 5055 4219 gi 62964 arylamine N-acetyltransferase 7 5055 4219 gi 62964 arylamine N-acetyltransferase 7 4141 4701 gi 682769 mccE gene product (Escherichia 10 6994 8742 gi 413951 ipa-27d gene product (Bacillus 10 6994 gi 413951 ipa-27d gene product (Bacillus 10 13 1377 2214 gi 490328 LORF F [unidentified]	1436965	oir S43914 S43914	54	37	1434
2 1090 1905 91467378 Unknown (Bacillus subtilis) 1 1322 663 911333761 Imposin II heavy chain [Naegler 13 12655 13059 pir[S00485 S004 gene 11-1 protein precursor 13 2030 3709 911303813 YqeW [Bacillus subtilis] 7 5055 4219 9162964 arylamine N-acetyltransferase 17 4141 4701 91682769 ImccE gene product [Escherichia 17 4141 4701 91682769 ImccE gene product [Escherichia 10 6994 8742 91413951 ipa-27d gene product [Bacillus 13 3377 2214 91490328 LORF F [unidentified]	11240016		54	38	513
1 1322 663 gi 1353761 myosin II heavy chain [Naegler 13 12655 13059 pir 500485 5004 gene 11-1 protein precursor 13 2030 3709 gi 1303813 YqeW [Bacillus subtilis] 7 5055 4219 gi 62964 arylamine N-acetyltransferase 7 4141 4701 gi 682769 mccE gene product [Escherichia 10 6994 8742 gi 413951 ipa-27d gene product [Bacillus 13 3377 2214 gi 490328 LORF F [unidentified]	467378 unknown (Bacillus		54	26	816
13 12655 13059 pir S00485 S004 gene 11-1 protein precursor 13 2030 3709 gi 1303813 YqeW Bacillus subtilis 17 5055 4219 gi 62964 arylamine N-acetyltransferase 17 4141 4701 gi 682769 mccE gene product Escherichia 10 6994 8742 gi 413951 ipa-27d gene product Bacillus 13 3377 2214 gi 490328 LORF F [unidentified]	1353761 myosin II heavy chain [Naegleria		54	22	1 099
3 2030 3709 gi 1303813 YqeW (Bacillus subtilis) 7 5055 4219 gi 62964 arylamine N-acetyltransferase (BA 1-290) [Ga 7 4141 4701 gi 682769 mccE gene product (Escherichia coli) 10 6994 8742 gi 413951 ipa-27d gene product (Bacillus subtilis) 3 3377 2214 gi 490328 LORF F [unidentified]	r S00485 S004 gene 11-1 protein precursor -	(fragments)	54	35	405
7 5055 4219 gi 62964 arylamine N-acetyltransferase (AA 1-290) (Ga arylamine N-acetyltransferase (EC 2.3.1.5)	11303813	_	54	34	1680
7 4141 4701 gi 882769 InccE gene product (Escherichia 10 6994 8742 gi 413951 ipa-27d gene product (Bacillus 3 3377 2214 gi 490328 LORF F [unidentified]	62964 arylamine N-acetyltransferase (AA 1-290) (Ga		54	33	837
10 6994 8742 gil413951 ipa-27d gene product [Bacillus 3 3377 2214 gil490328 LORF F [unidentified]	682769	_	54	31	561
3 3377 2214 gi 490328 LORF F	413951 ipa-27d gene product [Bacillus		54	28	1749
	1490328 LORF F		54	28	1164
341 4 3201 3614 gi 171959 myosin-like protein [Saccharomyces cerevisiae]	1171959		54	25	414

Table 2

729 936 921 489 906 1215 399 225 249 279 279 249 909 951 564 975 846 204 384 402 537 756 228 387 | % ident | length (nt) 41 35 45 36 40 25 56 36 34 34 31 29 32 20 38 32 45 30 24 33 33 32 42 54 - 8 sim |NADH dehydrogenase subunit 4 [Apis mellifera ligustica] pir|S52968|S52968 | NADH dehydrogenase chain 4 - honeybee itochondrion (SGC4) |similar to eukaryotic Na+/H+ exchangers [Escherichia coli] | sp|P32703|YJCE_ECOLI HYPOTHETICAL 60.5 KD PROTEIN IN SOXR-ACS NTERGENIC (2-hydroxyhepta-2,4-diene-1,7-dioate isomerase [Methanococcus jannaschii] |rodC (tag3) polypeptide (AA 1-746) [Bacillus subtilis] ir|S06049|S06049 | rodC protein - Bacillus subtilis p|P13485|TAGF_BACSU TECHOIC ACID fused envelope glycoprotein precursor [Friend spleen focus-forming irus] llow homology to P20 protein of Bacillus lichiniformis and bleomycin | acetyltransferase of Streptomyces verticillus (Bacillus subtilis) [hypothetical protein (SP:P10120) [Haemophilus influenzae] |pir|S30782|S307 |integrin homolog - yeast (Saccharomyces cerevisiae) |sensory protein kinase [Streptomyces hygroscopicus] No definition line found [Caenorhabditis elegans] |sp|P37348|YECE |HYPOTHETICAL PROTEIN IN ASPS 5'REGION (FRAGMENT) |ipa-37d qoxA gene product [Bacillus subtilis] |ornithine transcarbamylase [Homo sapiens] [hypothetical protein [Synechocystis sp.] |ORF YNL091w [Saccharomyces cerevisiae] |ORF YGR234w [Saccharomyces cerevisiae] |unknown [Corynebacterium glutamicum] liucA gene product [Escherichia coli] |product unknown (Bacillus subtilis) |sp|P02983|TCR S |TETRACYCLINE RESISTANCE PROTEIN. |ORF f343a [Escherichia coli] |ORF2 [Bacillus megaterium] | BIOSYNTHESIS PROTEIN F. |YqgZ [Bacillus subtilis] |ch-TOG [Homo sapiens] match gene name REGION (0549) acession |gi|1205015 |gi|1500558 1gi11303873 |gi|1063250 |gi|1301994 |gi|1001108 |gi|1323423 |gi|1045057 lgi|396400 |gi|474190 |gi|216267 |gi|304160 |gi|950197 gi|485090 gi|331906 |gi|219959 |gi|552446 1gi1987096 1gi1580870 |gi|537109 |gi|40100 1 5357 1351 1 1007 | 1942 1 2229 1 1522 1 4279 11011 (nt) 8854 1480 | 5301 11133 1 1441 1 1 1 1820 | 912 1 936 1 757 249 404 249 | 251 859 281 228 387 928 Contig |ORF | Start | ID |ID | (nt) | 1 2 | 2717 1 2 | 1137 1 9 1 7934 1 6262 1 1 1 2736 3305 1 1097 4351 1696 1356 11 11547 91 554 25 623 989 1 1 1 499 က 1314 | 1 | 3 - 2 - 2 -5 - 2 9 7 9 7 _ <u>س</u> -<u>-</u> --2450 346 378 444 463 505 558 580 603 617 622 623 408 502 551

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

3002 1 3561 1 3572 1	+	(1117)	acession		-		(311)
++	-	1 309	qi 44027	+	54	33	309
- ‡		1 464	 gi 151259	COA reductase (EC 1.1.1.88) [Pseudomonas mevalon	54	35	456
	- +	- +	- +	hydroxymethylglutaryl-CoA reductase (EC 1.1.1.88)	- +	- ‡	- +
	72	401	gi 450688	hsdM gene of EcoprrI gene product (Escherichia coli) pir 838437 838437 hsdM protein - Escherichia coli pir 809629 809629 hypothetical protein A - Escherichia coli (SUB 40-520)	54	36	330
3829 1	1 798	1 400	gi 1322245	mevalonate pyrophosphate decarboxylase [Rattus norvegicus]	54	29	399
3909 1		1 273	gi 29865	CENP-E [Homo sapiens]	54	30	273
3921 1	m 	1 209	pir S24325 S243	Iglucan 1,4-beta-glucosidase (EC 3.2.1.74) - Pseudomonas fluorescens subsp. cellulosa	54	34	207
4438 1	1 566	1 285	gi 1196657	unknown protein [Mycoplasma pneumoniae]	54	30	282
4459 1	8 -	1 272	gi 1046081	hypothetical protein (GB:D26185_10) [Mycoplasma genitalium]	54	38	270
4564 1	e -	1 221	gi 216267	ORF2 [Bacillus megaterium]	54	38	219
23 12	112538	110685	gi 474192	lucC gene product [Escherichia coli]	53	35	1854
23 14	114841	113579	gi 42029	ORF1 gene product [Escherichia coli]	53	32	1263
24 3	1 4440	1 3940	gi 1369947	1c2 gene product [Bacteriophage B1]	53	36 1	501
26 4	3818	1 4618	gi 1486247	unknown [Bacillus subtilis]	53	3.7	801
38 6	2856	1 3998	gi 405880	yeil [Escherichia coli]	53	40	1143
38 10	1 9380	9087	gi 1399954	thyroid sodium/iodide symporter NIS [Rattus norvegicus]	53	29	1575
56 10	112324	112100	pir A54592 A545	110k actin filament-associated protein - chicken	53	32	225
57 6	1 5047	1 4583	pir A00341 DEZP	alcohol dehydrogenase (EC 1.1.1.1) - fission yeast (Schizosaccharomyces pombe)	53	39	465
57 12	110515	8932	gi 1480429		53	30	1584
67 12	9496	110218	gi 1511555	[quinolone resistance norA protein protein [Methanococcus jannaschii]	53	31	723
69 - 3	1 3125	1 2382	gi 1087017	larabinogalactan-protein, AGP (Nicotiana alata, cell-suspension culture filtrate, Peptide, 461 aa]	53	30	744
79 1	e -	1031	gi 1523802	[glucanase [Anabaena variabilis]	53	32	1029
80 1	1 673	1 338	lgi1452428	ATPase 3 [Plasmodium falciparum]	53	36	336
88 4	1910	1 2524	gi 537034	ORF_0488 [Escherichia coli]	53	25	615
88 1 5	1 2467	1 3282	gi 537034	ORF_0488 [Escherichia coli]	53	29 1	816

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	+ Start (nt)	Stop (nt)	match acession	match gene name	# sim	% ident	length (nt)
92	80	1 5870	1 5505	qi 399598	amphotropic murine retrovirus receptor [Rattus norvegicus]	53	33	366
94		4 4417	3239	gi 173038	tropomyosin (TPM1) [Saccharomyces cerevisiae]	53	25	1179
66	5	1 4207	5433	sp P28246 BCR_E	HICYCLOMYCIN RESISTANCE PROTEIN (SULFONAMIDE RESISTANCE PROTEIN).	53	30	1227
120	3	1639	2262	gi 576655	ORF1 [Vibrio anguillarum]	53	35	624 1
1 120	111	1 7257	8897	gi 1524397	glycine betaine transporter OpuD [Bacillus subtilis]	53	33	1641
127	9 -	1 6893	5685	gi 1256630	putative [Bacillus subtilis]	53	32	1209
1 147	- 5	255	557	gi 581648	epiB gene product [Staphylococcus epidermidis]	53	34	303
158	+ ·	4705	4256	gi 151004	Imucoidy regulatory protein AlgR (Pseudomonas aeruginosa) pir A33802 A32802 regulatory protein algR - Pseudomonas aeruginosa sp P26275 ALGR_PSEAE Positive Alginote BlosynThEsis REGULATORY ROTEIN.	53	32	450
171	- 7	1 5717	5421	gi 1510669	hypothetical protein (GP:D64044_18) [Methanococcus jannaschii]	53	34	297
191	6 	13087	11483	gi 298085 	lacetoacetate decarboxylase [Clostridium acetobutylicum] pir B49346 B49346 butyrateacetoacetate CoA-transferase (EC .8.3.9) small chain - Clostridium acetobutylicum sp P33752 CTFA_CLOAB BUTYRATE-ACETOACETATE COA- TRANSFERASE SUBUNIT (EC 2.8.3.9) (COAT A)	53 33	31	1605
1 203	2	1 3763	1 4326	gi 143456	rpoE protein (ttg start codon) [Bacillus subtilis]	53	29	564
1 206	117	118204	118971	gi 304136 	acerylglutamate kinase (Bacillus stearothermophilus) sp 007905 ARGB_BACST ACETYLGLUTAMATE KINASE (BC 2.7.2.8) (NAG INASE) (AGK) (N-ACETYL-L-GLUTAMATE 5-PHOSFHOTRANSFERASE).	53	36	768
1 212	110	4021	4221	gi 9878		53	28	201
1 231	7 -	1580	1350	gi 537506	paramyosin [Dirofilaria immitis]	53	34	231
1 272	9	1 2719	3249	pir A33141 A331	hypothetical protein (gtfD 3' region) - Streptococcus mutans	53	34	531
1 308	8	1 927	1 2576	lgi1606292	ORF_0696 [Escherichia coli]	53	33	1650
320		1 5645	5884	gi 160596 	RNA polymerase III largest subunit (Plasmodium falciparum) sp P27625 RPC1_PLAFA DNA-DIRECTED RNA POLYMERASE III LARGEST UBUNIT (EC 2.7.7.6).	53	33	240
1 327		1 218	901	lgi1854601	unknown [Schizosaccharomyces pombe]	53	31	684
341	- 5	1 212	1 2500	lgi1633732	ORF1 [Campylobacter jejuni]	53	31	2289
351		1 763	1 383	sp P31675 YABM_	HYPOTHETICAL 42.7 KD PROTEIN IN TBPA-LEUD INTERGENIC REGION (ORF104).	53	32	381
433	7	1 5087	4731	lgi11001961	MHC class II analog [Staphylococcus aureus]	53	30	357
454		1240	086	pir A60328 A603	40K cell wall protein precursor (sr 5' region) - Streptococcus mutans (strain OM2175, serotype f)	53	27	261

| % ident | length (nt) # sim hsdM gene of EcoprrI gene product [Escherichia coli] pir|S38437|S38437 hsdM | protein - Escherichia coli pir|S09629|S09629 hypothetical protein A -|similar to unidentified ORF near 47 minutes [Escherichia coli] | sp|P31436|YICK_ECOLI HYPOTHETICAL 43.5 KD PROTEIN IN SELC-NLPA NTERGENIC ltoxic shock syndrome toxin-1 precursor (Staphylococcus aureus)
l pir/A24606|XCSAS1 toxic shock syndrome toxin-1 precursor - taphylococcus ir[S15765[S15765 hypothetical protein 1 (hlb 5' region) - aphylococcus 122.4% identity with Escherichia coli DNA-damage inducible protein ...; ORF 1 (AA 1 - 121) (1 is 2nd base in codon) [Staphylococcus aureus] S. aureus - Putative coding regions of novel proteins similar to known proteins |putative transcriptional regulator [Bacillus stearothermophilus] orfA2; orfA2 forms an operon with orfA1 (Listeria monocytogenes) |pilin biogenesis protein [Haemophilus influenzae] No definition line found [Caenorhabditis elegans] transmembrane protein (Lactococcus lactis) |lipase-like enzyme [Alcaligenes eutrophus] !myosin heavy chain [Entamoeba histolytica] Ihypothetical protein [Synechocystis sp.] ORF 3; putative [Bacillus subtilis] |unknown [Schizosaccharomyces pombe] |transposase [Burkholderia cepacia] |SapA protein [Escherichia coli] Escherichia coli (SUB 40-520) Hmp (Vibrio parahaemolyticus) putative [Bacillus subtilis] |res gene [Bacteriophage Pl} |rat GCP360 [Rattus rattus] |YqiX [Bacillus subtilis] |EF [Streptococcus suis] aureus (fragment) match gene name REGION. acession |gi|1480429 |gi|11146243 |gi|1279400 |gi|1303949 lgi11477486 |gi|1204551 |gi|1044936 |gi|1001774 |gi|695278 |gi|516826 match 1gi1507738 |gi|153123 gi|687600 |gi|290508 |gi|142441 1gi | 305080 gi|308852 |gi|450688 | 5 | 3049 | 3441 |gi|868224 |gi|298032 |gi|46587 |gi|15140 (nt) Start | Stop 1 590 1 504 1 434 1 184 1 397 (nt) 1 1 1004 1 3957 ო — ~ ~ m Contig |ORF | -ΩI -_ --

 Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	I JORF	Start (nt)	Stop (nt)	match acession	match gene name	e sim	% ident	l length (nt)
15		1 2205	1 2369	gi 215966	G41 protein (gtg start codon) [Bacteriophage T4]	52	34	165
19	e -	1 2429	3808	gi 1205379	UDP-murnac-pentapeptide synthetase [Haemophilus influenzae]	52	31	1380
24		6920	3462	gi 579124 	predicted 86,4kd protein; 52Kd observed [Mycobacteriophage 15] pir 530971 530971 gene 26 protein - Mycobacterium phage L5 sp Q05233 VG26_BPML5 MINOR TAIL PROTEIN GP26. (SUB 2-837)	52	32	3459
1 37		3015	3935	gi 1500543		52	1 25	921
1 38	113	8795	1 9703	 gi 46851		52	29	1 606
1 44	116	110617	111066	gi 42012	moaE gene product [Escherichia coli]	52	36	450
1 46		e 	521	gi 1040957		52	25	519
51	110	5531	1 6280	gi 388269		52	32	750
29	~	3968	1 2826	gi 181949 	endothelial differentiation protein (edg-1) [Homo sapiens] pir[A35300[A35300 G protein-coupled receptor edg-1 - human sp P21453 EDG1_HUMAN PROBABLE G PROTEIN-COUPLED RECEPTOR EDG-1.	52	23	1143
1 57	— ت	1 4850	4173	gi 304153		52	27	678
62	- 2	3364	1 2870	gi 1072399		52	25	495
1 62	9 -	4445	3651	gi 46485	NADH dehydrogenase [Synechococcus PCC7942]	52	27	1 267
1 67	114	111355	112962	(gi 1511365	Iglutamate synthase (NADPH), subunit alpha [Methanococcus jannaschii]	52	30	1608
1 67	21	116935	118158	gi 1204393	hypothetical protein (SP:P31122) [Haemophilus influenzae]	52	25	1224
1 70	4	1 2185	1997	gi 7227 		52	36	189
96	110	110005	110664	gi 1408485	[B65G gene product [Bacillus subtilis]	52	26	099
103	- 5	3986	1 3351	gi 1009368	Respiratory nitrate reductase (Bacillus subtilis)	52	42	636
1 109	- 3	4102	1 3350	gi 699274	ImbE gene product [Mycobacterium leprae]	52	39	753
1 109	119	115732	117300	gi 1526981	lamino acid permease YeeF like protein [Salmonella typhimurium]	52	30	1569
121	3	1412	981	gi 732931		52	32	432
125	3	865	1680	(gi/1296975		52	38	816
130	5	629	1807	gi 1256634 	[25.8% identity over 120 aa with the Synenococcus sp. MpeV protein; putative [Bacillus subtilis]	52	36	1149
149	. –	11164	1 583	gi 1225943	PBSX terminase [Bacillus subtilis]	52	33	582
149	114	1 4687	4415	gi 1510368	[M. jannaschii predicted coding region MJ0272 [Methanococcus jannaschii]	52	35	273

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)	match acession	match gene name	# sim	% ident	length (nt)
167	-	216	1001	gi 146025	cell division protein [Escherichia coli]	52	43	1986
188		120	1256	gi 474915 	orf 337; translated orf similarity to SM: BCR ECOLI bicyclomycin esistance protein of Escherichia coli (Coxiella burnetii) pir S44207 S44207 hypothetical protein 337 - Coxiella burnetii (SUB -338)	52	26	1137
1 195	6	9161	1 8760	qi 3028	mitochondrial outer membrane 72K protein (Neurospora crassa) r A36682 A36682 72K mitochondrial outer membrane protein - rospora crassa	52	25	402 +
1 200	<u>س</u>	1 2065	1 2607	gi 142439	ATP-dependent nuclease [Bacillus subtilis]	52	35	543
1 203	4	1 2776	3684	gi 1303698	BitD (Bacillus subtilis)	52	25	+ 606
1 227	œ 	1 5250	1 5651	gi 305080	myosin heavy chain [Entamoeba histolytica]	52	24	402
1 242	1	21	1424	gi 1060877	EmrY [Escherichia coli]	52	32	1404
1 249		4526	1 4753	pir C37222 C372	pir C37222 C372 cytochrome P450 lA1, hepatic - dog (fragment)	52	23	228
255		1 2107	1055	gi 143290	penicillin-binding protein [Bacillus subtilis]	52	28	1053
276		1 3963	3664	gi 1001610	hypothetical protein (Synechocystis sp.)	52	30	300
276	œ —	1 4456	4055	gi 416235	orf L3 [Mycoplasma capricolum]	52	26	402
289	- 2	1856	1449	gi 150900	GTP phosphohydrolase [Proteus vulgaris]	52	34	408
325			279	gi 1204874 	polypeptide deformylase (formylmethionine deformylase) [Haemophilus influenzae]	52	33	279
340		2017	1010	gi 1215695 	peptide transport system protein SapF homolog; SapF homolog [Mycoplasma pneumoniae]	52	33	1008
375		340	1 1878	gi 467446	similar to SpoVB [Bacillus subtilis]	52	28	1539
424	4	4104	3262	gi 1478239	unknown (Mycobacterium tuberculosis)	52	34	843
430		3	575	pir A42606 A426	orfA 5' to orf405 - Saccharopolyspora erythraea (fragment)	52	28	573
644	4	4728	3712	gi 1408494	homologous to penicillin acylase [Bacillus subtilis]	52	31	1017
465		1802	903	gi 143331 	alkaline phosphatase regulatory protein (Bacillus subtilis) pir A27650 A27650 regulatory protein phoR - Bacillus subtilis sp P23545 PHOR_BACSU ALKALINE PHOSPHATASE SYNTHESIS SENSOR PROTEIN HOR (EC 2.7.3).	52	36	006
469	2	4705	4169	gi 755152 	highly hydrophobic integral membrane protein [Bacillus subtilis] sp P42953 TAGG_BACSU TEICHOIC ACID TRANSLOCATION PERWEASE PROTEIN AGG.	52	32	537
1 495		1262	1 633	gi 1204607	transcription activator [Haemophilus influenzae]	52	25	630
505	_	6004	1 5762	gi 142440	ATP-dependent nuclease (Bacillus subtilis)	52	28	243
1	! !				+ + + + + + + + + + + + + + + + + + + +	1 1 1 1 1 1 1 1 1		

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Bacteriophage phi-11 int gene activator [Staphylococou	Contig	108F ID	Start (nt)	Stop (nt)	match acession	match gene name	s sim	% ident	length (nt)
1 1 336 gil1215693 putative orf; GT9_Orf434 [Mycoplasma pneumoniae] 1 1 336 gil551648 lep1B gene product [Staphylococcus ejidermidis] 1 1 336 gil551648 lep1B gene product [Staphylococcus ejidermidis] 1 1 324 gil1279769 [Pdhc [Hethanobacterium thermoformidicum] 1 324 gil126131 [Pdhc [Hethanobacterium thermoformidicum] 1 324 gil126131 [Pdhc [Hethanobacterium thermoformidicum] [Pdhc [Hethanobacter [Hethanobacter Phylori] [Pdhc [Hetha			1162	1614		Bacteriophage phi-11 int gene activator [Staphylococcus acteriophage phi 11]	52	35	453
1 1 336 gil581648 lepiB gene product [Staphylococcus epidermidis] 1 848 446 gil129769 Frbhc [Methanobacterium thermoformicicum] 1 1 324 gil14439 MAP-dependent nuclease [Bacillus subtilis] 1 1 324 gil126322 comm Ostra Garacteriophage SPO2 produced Garacteriophage Garacte	543		444	1295	gi 1215693	orf; GT9_orf434 [Mycoplasma	52	25	852
1 848 426 gil1279769 Fedro [Methanobacterium thermoformicicum] 1 130 330 gil14439 IAPP-dependent nuclease [Bacillus subtilis] 1 1324 gil1289262 Commc ORF3 [Bacillus subtilis] 1 1 1 1 1 1 1 1 1	586		1	336	gi 581648	gene product [Staphylococcus	52	36'	336
1 691 330 91 142439 ATP-dependent nuclease [Bacillus subtilis] 1 691 347 91 289262 comE ORF3 [Bacillus subtilis] 1 324 91 216512 SP02 DNA POLYMERASE (as 1-648) [Bacteriophage SP02] 1 566 285 91 1256136 NDA POLYMERASE (as 1-648) [Bacteriophage SP02] 1 588 295 91 298032 EF [Streptococcus suis] 1 588 295 91 298032 EF [Streptococcus suis] 1 588 295 91 298032 EF [Streptococcus suis] 1 560 91 218040 BBA [Bacillus lichenifornis] 1 478 260 91 218040 BBA [Bacillus lichenifornis] 1 478 240 91 1205919 Influenced and anihobutrylc acid transport in thuman lichenifornis 1 4 4 4 4 4 4 4 4 4	1 773	-	848	426	gi 1279769		52	30	423
1 691 347 991289262	1120	-	100	330		ATP-dependent nuclease [Bacillus subtilis]	52	35	231
1 1 324 94 216151 IDNA polymerase (gene L; trg start codon) [Bacteriophage PO2] 1 566 285 94 1256136 [YbbG [Bacillus subtilis]] 1 566 285 94 1256136 [YbbG [Bacillus subtilis]] 1 568 295 94 1236136 [YbbG [Bacillus subtilis]] 1 588 295 94 1236136 [YbbG [Bacillus subtilis]] 1 588 295 94 1236025 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 562 94 1236025 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 562 94 1236025 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 562 94 1236025 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 562 94 1236046 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 560 94 1236046 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 590 401 94 1236046 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 549 94 140003 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 549 94 1530936 [Phypothetical 64.7-EDa protein [Bacillus subtilis]] 1 575 94 1955077 [Phypothetical 64]1000000000000000000000000000000000000	1614	-	691	347	lgi1289262	comE ORF3 [Bacillus subtilis]	52	28	345
1 566 285 gil1256136	2495			324	gi 216151 	DNA polymerase (gene L; ttg start codon) [Bacteriophage SP02] gil579197 SP02 DNA polymerase (aa 1-648) [Bacteriophage SP02] pir A21498 DJBPS2 DNA- directed DNA polymerase (EC 2.7.7.7) - phage P02	52	34	324
1 577 320 91 41713 hisA ORF (AA 1-245) [Escherichia coli] 1 588 295 91 298032 EF [Streptcococus suis] 1 612 307 91 49925 hypothetical 64.7-kBa protein [Bacillus subtilis] 1 478 260 91 1218040 BAA [Bacillus licheniformis] 2 613 347 91 1205919 Ma+ and Cl- dependent gamma-aminobutryic acid transpor influenzae 1 3 389 91 40003 2-0x06ullurarate dehydrogenase (WADP+) [Bacillus subtilis] 2-0x06ullurarate dehydrogenase (WADP+) [Bacillus subtilis] 1 1 249 91 159388	1 2931	-	566	285	gi 1256136	[Bacillus	52	30	282
1 588 295 gi1298032 EF [Streptococcus suis]	1 2943		577	320	gi 41713	(AA	52	35	258
1 612 307 gil899025 hypothetical 64.7-kDa protein [Bacillus subtilis] 1 478 260 gil1218040 BAA [Bacillus licheniformis] 2 613 347 gil854064 1087 [Human herpesvirus 6] 1 90 401 gil1205919 Na+ and Cl- dependent gamma-aminobutryic acid transpor 1 3 389 gil40003 c2coxocuprante Dehydrogenase (NADP+) [Bacillus subtilis] 1 1 1 249 gil1205919 No definition line found [Escherichia coli) DEHYDROGENASE] DEHYDROGENASE EL COMPONENT (EC 2.4.2) DEHYDROGENASE DEHYDROGENASE CAPONENT (EC 2.4.2) DEHYDROGENASE] No definition line found [Escherichia coli) 1 3 1212 gil180507 No definition line found [Escherichia coli) 1 3 575 gil180507 No definition line subtilis] 1 3 575 gil180507 No definition subtilis 1 3 596 gil130993 YqiN [Bacillus subtilis] 1 1 1 1 1 1 1 1 1	2993		588	295	lgi1298032	[EF [Streptococcus suis]	52	34	294
1 478 260 gil1218040 BAA [Bacillus licheniformis]	3667	_	612	307		64.7-kDa protein [Bacillus	52	36	306
1 90 401 1911205919 INA+ and CI- dependent gamma-aminobutryic acid transpor influenzae] 1 30 401 1911205919 Influenzae] Influenzae] 1 3 389 191140003 Oxoglutarate dehydrogenase (NADP+) [Bacillus subtilis] 2-OXOGLUTARATE DEHYDROGENASE El COMPONENT (EC 2.4.2) DEHYDROGENASE) 1 249 191159388 Ornithine decarboxylase [Leishmania donovani] 2-OXOGLUTARATE DEHYDROGENASE El COMPONENT (EC 2.4.2) DEHYDROGENASE) Influence continuental denovani 2-OXOGLUTARATE DEHYDROGENASE El COMPONENT (EC 2.4.2) 1 3 212 1911955077 No definition line found (Escherichia coli) 1 3 275 1911895747 Putative cel operon regulator (Bacillus subtilis) 2 2 2 2 2 2 2 2 2	3944		478	260	gi 1218040	BAA [Bacillus licheniformis]	52	36	219
1 90 401 gi 1205919 Na+ and Cl- dependent gamma-aminobutryic acid transpor 3 389	3954	-	613	347	lgi1854064	U87 [Human herpesvirus 6]	52	50	267
1 3 389 igi140003 cxcglutarate dehydrogenase (NADP+) [Bacillus subtilis] 2-0x0GLUTARATE DEHYDROGENASE EI COMPONENT (EC 2.4.2) DEHYDROGENASE EI COMPONENT (EC 2.4.2) EHYDROGENASE EI COMPONENT (EC 2.4.2) ET 1 1 1 1 1 249 igi1159388	3986		06	401		Na+ and Cl- dependent gamma-aminobutryic acid transporter (Haemophilus influenzae]	52	33	312
1	4002		— — — -	389	gi 40003 	(NADP+) [Bacillus SE El COMPONENT (F	52	42	387
1 438 220 gi 409795 1 3 212 gi 965077 4 2479 3276 gi 11510962 9 5501 5966 gi 11510962 9 5501 5966 gi 11510460 17 11042 11305 gi 142011 11 6453 6731 gi 1495471	4020		1	249	gi 159388	ornithine decarboxylase [Leishmania donovani]	52	47	249
1 3 212	1 4098	- 1	438	220	gi 409795	found (Escherichia	52	32	219
1 3 575 g1 895747 4 2479 3276 g1 1510962 9 5301 5966 g1 1303933 3 1516 1283 g1 1519460 17 11042 11305 g1 42011	4248		3	212			52	40	210
4 2479 3276 gi 1510962 9 5301 5966 [gi 1303933 3 1516 1283 gi 1519460 17 11042 11305 gi 42011 11 6453 6731 [gi 495471	7	-	3	575	lgi1895747	operon regulator [Bacillus	51	28	573
9 5301 5966 (4) 1303933 3 1516 1283 (4) 11519460 17 11042 11305 (4) 42011 11 6453 6731 (4) 495471	21	-	2479	3276	gi 1510962	indole-3-glycerol phosphate synthase [Methanococcus jannaschii]	51	32	1 867
3 1516 1283	22		5301	5966	lgi11303933	YqiN [Bacillus subtilis]	51	25	1 999
17 11042 11305 91 42011 11 6453 6731 91 495471	43		1516	1283	gi 1519460		51	31	234
11 6453 6731 gi 495471	i		:	11305	gi 42011	moaD gene product [Escherichia coli]	51	35	264
		-	6453	6731	gi 495471	vacuolating toxin [Helicobacter pylori]	51	37	279

Table 2

| % ident | length (nt) | % sim EIIA domain of PTS-dependent Gat transport and phosphorylation Escherichia rodD (gtaA) polypeptide (AA 1-673) [Bacillus subtilis] pir|S06048|S06048| probable rodD protein - Bacillus subtilis sp|P13484|TAGE BACSU PROBABLE | POLY(GLYCEROL-PHOSPHATE) LPHA-GLUCOSYLTRANSFERASE (EC 2.4.1.52) (TECHOIC M. jannaschii predicted coding region MJ0938 [Methanococcus jannaschii] 125% identity to the E.coli regulatory protein MprA; putative [Bacillus |pir|A45605|A456 |mature-parasite-infected erythrocyte surface antigen MESA - Plasmodium (H. influenzae predicted coding region HI0056 [Haemophilus influenzae] |pir|S05330|S053 |maltose-binding protein precursor - Enterobacter aerogenes (hypothetical protein (SP:P32662) (Haemophilus influenzae) |hypothetical protein (SP:P33918) [Haemophilus influenzae] proliferating cell nuclear antigen [Styela clava] |melvalonate kinase [Methanococcus jannaschii] Inuclear protein [Drosophila melanogaster] |alanine dehydrogenase (Bacillus subtilis) |MunI regulatory protein [Mycoplasma sp.] |ORF YDL085w [Saccharomyces cerevisiae] [Asparaginase [Bacillus licheniformis] [TFIID subunit TAFII55 [Homo sapiens] |unknown [Mycobacterium tuberculosis] [B1496 C3 206 [Mycobacterium leprae] ACID BIOSYNTHESIS ROTEIN E). |orf1 [Mycoplasma capricolum] |unknown [Bacillus subtilis] [bmrU [Bacillus subtilis] |yehU [Escherichia coli] |EF [Streptococcus suis] match gene name subtilis] coli] acession |gi|1256652 |gi|1161242 |gi|1204815 |gi|1431110 |gi|1205484 |gi|1510977 |gi|1204314 1gi11237044 1gi1466886 match |gi|508173 |gi|467359 1gi1298032 |gi|642795 | 1428 |gi|580920 |gi|405857 1gi1435098 |gi|431929 |gi|409286 |gi|299163 1gi149272 1gi18204 4 | 3357 | 4040 1 6693 1 2352 | 9647 | 10477 1 7356 (nt) 1 8967 Start | (nt) 1 4 1 4637 1 1 2852 1 6007 1 7331 1 707 ლ — Contig | ORF | ID | - _ - 5 <u>~</u> -_ œ — ญ

Table

S. aureus - Putative coding regions of novel proteins similar to known proteins

					+	++		+
Contig ID	IORF	Start (nt)	Stop (nt)	match acession	match gene name	mis %	% ident	length (
1 257	4	3540	1 3373	gi 1204579	H. influenzae predicted coding region HI0326 [Haemophilus influenzae]	51	22	168
258	e	2397	1609	gi 160299	glutamic acid-rich protein (Plasmodium falciparum) pir1A545141A54514 glutamic acid-rich protein precursor - Plasmodium alciparum	51	34	789
265		2419	1 3591	gi 580841	F1 [Bacillus subtilis]	51	32	1173
298	1 2	518	1 748	gi(1336162		51	34	231
316	6	5817	1 7049	gi 413953		51	39	1233
332	2	3775	1 2057	gi 1209012	mutS [Thermus aquaticus thermophilus]	51	26	1719
364	4	3816	4991	gi 528991	unknown [Bacillus subtilis]	51	32	1176
440	7	448	684	·	Exansferase (GAL10) (AA 1 - 687) [Kluyveromyces lactis] r 501407 XUVKG UDPglucose 4-epimerase (EC 5.1.3.2) - yeast uyveromyces marxianus var. lactis)	51	32	237
495	2	1353	7,111	gi 297861		51	41	1771
495	3	1 2287	1718	gi 1513317		51	25	570
206		840	421	gi 455320	cII protein (Bacteriophage P4]	51	33	420
009		1474	983	gi 587532 	orf, len: 201, CAI: 0.16 [Saccharomyces cerevisiae] pir 548818 548818 hypothetical protein - yeast (Saccharomyces erevisiae)	51	30	492
607	3	479	934	gi 1511524	hypothetical protein (SP:P37002) [Methanococcus jannaschii]	51	40	456
989	2	127	009 1	lgi 493017	[endocarditis specific antigen [Enterococcus faecalis]	51	30	474
726	1	33	1 230	gi 1353851	unknown [Prochlorococcus marinus]	51	45	198
861		176	1 652	gi 410145	dehydroquinate dehydratase [Bacillus subtilis]	51	34	477
869	-	782	393	gi 40100 	rodC (tag3) polypeptide (AA 1-746) [Bacillus subtilis] ir S06049 S06049	51	23	390
1003		642	1 322	lgi11279707	hypothetical phosphoglycerate mutase [Saccharomyces cerevisiae]	51	39	321
1046	1 2	998	624	gi 510257	glycosyltransferase [Escherichia coli]	51	29	243
1467		702	352	gi 1511175	[M. jannaschii predicted coding region MJ1177 [Methanococcus jannaschii]	51	32	351
2558		457	1 230	sp P10582 DPOM_	DNA POLYMERASE (EC 2.7.7.7) (S-1 DNA ORF 3).	51	26	228
3003		677	1 399	gi 809543	CbrC protein [Erwinia chrysanthemi]	1 51	27	381
3604		1	1 399	pirlJC42101JC42	irlJC4210 JC42 3-hydroxyacyl-CoA dehydrogenase (EC 1.1.1.35) - mouse	51	37	399
3732	- 1	2	316	gi 145906	acyl-CoA synthetase [Escherichia coli]	51	33	315
						-		

Table 2

| % ident | length | (nt) 8 sim l coded for by C. elegans cDNA cm01e7; Similar to hydroxymethylglutaryl-CoA synthase [Caenorhabditis elegans] IORF_f167; end overlaps end of o100 by 14 bases; start overlaps f174, ther |GRAIL score: null; cap site and late promoter motifs present pstream; ribosomal protein L12 (AA 1-179) [Escherichia coli] ir|S04776|XXECPL | peptide N-acetyltransferase rimL (EC 2.3.1.-) - cherichia coli Similar to dihydroflavonol-4-reductase (maize, petunia, tomato) |Similar to dihydroflavonol-4-reductase (maize, petunia, tomato) putative [Autographa californica nuclear polyhedrosis irus] carbamoyl phosphate synthetase II [Plasmodium falciparum] [homologous to penicillin acylase [Bacillus subtilis] |xylitol dehydrogenase [unidentified hemiascomycete] |LmrP integral membrane protein [Lactococcus lactis] putative cel operon regulator [Bacillus subtilis] circumsporozoite protein [Plasmodium reichenowi] |semaphorin III family homolog [Homo sapiens] |structural protein [Bacteriophage Tuc2009] hypothetical protein [Synechocystis sp.] |surfactin synthetase [Bacillus subtilis] lipa-42d gene product [Bacillus subtilis] ORF174 gene product [Porphyra purpurea] |ORF YGR103w [Saccharomyces cerevisiae] unknown [Mycobacterium tuberculosis] [FemA protein [Staphylococcus aureus] | starts possible [Escherichia coli] |putative (Bacillus subtilis) |putative [Bacillus subtilis] [Caenorhabditis elegans] | [Caenorhabditis elegans] |Yqjo [Bacillus subtilis] | match gene name acession |gi|1408494 |gi|1458280 |gi|1208451 |gi|1072179 |gi|1061351 |gi|1146207 |gi|1323159 |gi|1303966 |gi|1072179 |gi|1276658 1gi11403441 |gi|1052754 |gi|216346 |gi|559160 |gi|413966 |gi|153015 1gi | 606096 |gi|640922 1gi 1476024 |gi|895747 |gi|496280 |gi|143725 |gi|160229 |gi|42749 1105 | 2976 1 9496 1 4824 113 (15290 (15841 Start | Stop (nt) | (nt) 1 4859 1 2759 1 5096 1 274 1 9623 1 626 14 | 3576 14 | 1926 1 1 1 367 ო — Contig |ORF | ID |ID | ----_ - ر <u>-</u> _ Ξ -

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	- C		Ston			+		++
QI			(nt)	acession	מתרכנו מנודים			(nt)
211	- 4	1523	1927	gi 410131	ORFX7 [Bacillus subtilis]	05 1	29	405
214	4	2411	3295	sp P37348 YECE_	HYPOTHETICAL PROTEIN IN ASPS 5'REGION (FRAGMENT).	20	37	885
228		1 5068	1 4406	gi 313580 	envelope protein [Human immunodeficiency virus type 1] pir[835835 835 envelope protein - human immunodeficiency virus ype 1 (fragment) [SUB 1- 77]	20	35	663
272		3048	1723	lgi11408485	1865G gene product (Bacillus subtilis)	20	22	1326
1 273	7	1616	984	gi 984186	phosphoglycerate mutase (Saccharomyces cerevisiae)	20	28	633
328	7	1 2507	1605	gi 148896	lipoprotein [Haemophilus influenzae]	20	26	903
1 332	4	1 5469	3802	gi 1526547	DNA polymerase family X [Thermus aquaticus]	20	27	1668
342	2	3473	3931	gi 456562	G-box binding factor [Dictyostelium discoideum]	50	35	459
352		1 1478	741	gi 288301	ORF2 gene product [Bacillus megaterium]	20	29	738
408	1 7	1 5299	5523	gi 11665	ORF2136 [Marchantia polymorpha]	20	27	225
420	<u>س</u>	1 650	1825	gi 757842	UDP-sugar hydrolase [Escherichia coli]	20	30	1176
1 464			591	gi 487282	Na+ -ATPase subunit J [Enterococcus hirae]	20	29	591
472	- 5	1418	1 864	gi 551875	BglR [Lactococcus lactis]	20	23	555
1 520		1 23	541	gi 567036	CapE [Staphylococcus aureus]	50	27	519
529		9	410	gi 1256652 	[25% identity to the E.coli regulatory protein MprA; putative [Bacillus subtilis]	20	34	405
534	ιn	1 7726	6029	gi 295671 	selected as a weak suppressor of a mutant of the subunit AC40 of DNA ependant RNA polymerase I and III {Saccharomyces cerevisiae}	50	18	1668
647		1 2990	1497	gi 405568 	Tral protein shares sequence similarity with a family of opoisomerases [Plasmid pSK41]	20	31	1494
664	. — -	1133	117	gi 410007	leukocidin F component [Staphylococcus aureus, MRSA No. 4, Peptide, 23 aa]	50	32	423
678			627	gi 298032	[EF [Streptococcus suis]	50	29	627
755	e 	947	1171	gi 150572 	Cytochrome cl precursor (EC 1.10.2.2) (Paracoccus denitrificans) gi145465 cytochrome cl (AA 1-450) [Paracoccus denitrificans) pir1C29413 c29413 ubiquinolcytochrome-c reductase (EC 1.10.2.2) ytochrome cl precursor - Paracoccus denitrificans sp[P13627(CY1	20	37	225
827		1363	683	gi 142020	heterocyst differentiation protein [Anabaena sp.]	20	21	681
892		m 	752	gi 1408485	B65G gene product [Bacillus subtilis]	20	27	750
910	7	1 438	1 887	gi 1204727	ltyrosine-specific transport protein [Haemophilus influenzae]	20	25	450
1	+	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	! ! ! ! !		+ 1 1 1 1 1 1 1 1 1 1			

Table 7

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	mis &	m % ident	l length (nt)
933		524	1 760	gi 1205451	cell division inhibitor [Haemophilus influenzae]	- 50	0 32	237
973		424	1 236	lgi1886947	orf3 gene product [Saccharomyces cerevisiae]		50 40	189
1009		653	1 429	gi 153727	M protein [group G streptococcus]	1 20	0 28	1 225
1027		511	1 257	gi 413934	lipa-10r gene product (Bacillus subtilis)	50	0 25	1 255
1153	1 2	1 556	1 326	gi 773676	InccA (Alcaligenes xylosoxydans)	200	36	231
1222		1 798	1 400	gi 1408485	B65G gene product [Bacillus subtilis]	1 50	0 21	399
1 1350		1 692	399	gi 289272	[ferrichrome-binding protein [Bacillus subtilis]	5	50 32	1 294
2945		366	184	gi 171704	hexaprenyl pyrophosphate synthetase (COQ1) [Saccharomyces erevisiae]	5	50 34	183
1 2968	1 2	1604	1 804	gi 397526	clumping factor [Staphylococcus aureus]	2	50 33	1 801
1 2998	2 -	1 657	394	gil495696	F54E7.3 gene product [Caenorhabditis elegans]	S	50 40	1 264
3046	1 2	905	306	pir S13819 S138	acyl carrier protein - Anabaena variabilis (fragment)	20	0 32	201
3063		1 547	1 275	gi 474190	lucA gene product [Escherichia coli]	20	0 1 29	1 273
3174		е —	146	gi 151900	alcohol dehydrogenase [Rhodobacter sphaeroides]	1 50	0 31	1 144
3792		625	314	gi 1001423	hypothetical protein [Synechocystis sp.]	20	0 35	312
3800		5	1 262	gi 144733 	NAD-dependent beta-hydroxybutyryl coenzyme A dehydrogenase Clostridium acetobutylicum	20	0 28	261
3946		1 373	188	gi 576765	cytochrome b [Myrmecia pilosula]		50 38	186
3984		1 578	1 291	sp(P37348 YECE_	HYPOTHETICAL PROTEIN IN ASPS 5'REGION (FRAGMENT).	20	0 37	1 288
37	110	8250	1 7885	gi 1204367	hypothetical protein (GB:U14003_278) [Haemophilus influenzae]	4	49 30	366
1 46	116	113802	114848	gi 466860	acd; B1308_F1_34 [Mycobacterium leprae]	4	49 24	1047
65	5	1 2267	3601	gi 606304	ORF_0462 [Escherichia coli]	49	9 27	1335
112	118	17884	118615	gi 559502	ND4 protein (AA 1 - 409) (Caenorhabditis elegans)	49	9 1 25	732
138	6	1 6973	1 7902	gi 303953	esterase [Acinetobacter calcoaceticus]	49	9 29	930
1 217	9	1 4401	1 5138	gi 496254	fibronectin/fibrinogen-binding protein [Streptococcus pyogenes]	4	49 31	738
1 220	112	111803	112657	gi 397526	[clumping factor [Staphylococcus aureus]	49	9 (31	855
1 228	4	1 1842	2492	pir \$23692 \$236	hypothetical protein 9 - Plasmodium falciparum	49	9 24	651
1 268	1	5016	1 2614	gi 143047	ORFB [Bacillus subtilis]	49	9 1 56	2403
: : : : : : :	! !!!!	 	+		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	1 1 1 1		

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	TID	Start (nt)	Stop (nt)	match acession	match gene name	# 8 sim	* ident	length (nt)
271	- 5	1164	1373	gi 1001257	hypothetical protein [Synechocystis sp.]	49	38	210
300	3	1 4340	3180	gi 1510796	hypothetical protein (GP:X91006_2) [Methanococcus jannaschii]	49	26	1161
381		2281	1142	 gi 396301 	matches PS00041: Bacterial regulatory proteins, araC family ignature [Escherichia coli]	4	29	1140
1 466		. 3	947	gi 1303863	VggP [Bacillus subtilis]	49	26	945
999		379	191	gi 633112	ORFI [Streptococcus sobrinus]	49	29	189
1 670	1 2	1 403	1014	gi 1122758	unknown [Bacillus subtilis]	49	32	612
607		1433	795	gi 143830	xpaC [Bacillus subtilis]	49	29	639
831		943	473	gi 401786	phosphomannomutase [Mycoplasma pirum]	49	29	471
1 1052		1 422	213	gi 1303799	YqeN [Bacillus subtilis]	49	21	210
1 1800		342	172	gi 216300 	peptidoglycan synthesis enzyme (Bacillus subtilis) sp P37585 MURG_BACSU MURG PROTEIN UPD-N-ACETYLGLUCOSAMINEN-ACETYLMURAMYL- PENTAPEPTIDE) FYROFHOSPHORYL-UNDECAPRENOL N-ACETYLGLUCOSAMINE RANSFERASE).	49	28	171
1 2430		2	376	sp P27434 YFGA_	HYPOTHETICAL 36.2 KD PROTEIN IN NDK-GCPE INTERGENIC REGION.	49	26	375
3096		542	1 273	gi 516360	surfactin synthetase [Bacillus subtilis]	49	25	270
32	- 4	3771	3100	gi 1217963	hepatocyte nuclear factor 4 gamma (HNF4gamma) [Homo sapiens]	48	36	672
38		1	609	gi 1205790	H. influenzae predicted coding region HI1555 [Haemophilus influenzae]	48	28	609
45	9	5021	6427	gi 1524267	unknown [Mycobacterium tuberculosis]	48	20	1407
65	114	116346	131096	gi 1197336	Lmp3 protein (Mycoplasma hominis)	48	28	14751
61	. —	3	809	gi 1511555	[quinolone resistance norA protein protein [Methanococcus jannaschii]	48	30	909
61	. — ·	3311	3646	gi 1303893	YqhL [Bacillus subtilis]	48	29	336
114		86	415	gi 671708 	su(s) homolog; similar to Drosophila melanogaster suppressor of able (su(s)) protein, Swiss-Prot Accession Number P22293 Drosophila virilis]	48	25	318
121		1131	610	gi 1314584	unknown {Sphingomonas S88}	48	29	522
1 136	-	2014	1280	gi 1205968	H. influenzae predicted coding region H11738 [Haemophilus influenzae]	48	23	735
171	110	8220	9557	gi 1208454	hypothetical protein [Synechocystis sp.]	48	34	1338
175		3625	1814	gi 396400 	similar to eukaryotic Na+/H+ exchangers [Escherichia coli] sp P32703 YJCE_ECOLI HYPOTHETICAL 60.5 KD PROTEIN IN SOXR-ACS NTERGENIC REGION (0549).	48	29	1812
1 194		2	385	gi 1510493	[M. jannaschii predicted coding region MJ0419 [Methanococcus jannaschii]	48	25	384
	-			-				

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

-)	(nt)	acession		_		(nt)
- /61	1 0	901	452	gi 1045714		48	25	450
203		·	396	gi 940288		48	60	396
204	1 - 1	1363	869	gi 529202	No definition line found [Caenorhabditis elegans]	48	25	999
206 12	20 134	34815	127760	911511490		48	27	7056
212		2	166	gi 295899	Inucleolin [Xenopus laevis]	48	34	165
220 1	110 112	12652	111426	gi 44073		48 -	23	1227
243	9 1 9	6450	5491	gi 1184118	mevalonate kinase [Methanobacterium thermoautotrophicum]	48	30	096
264	4 1 5	5434	3308	gi 1015903	ORF YJR151c [Saccharomyces cerevisiae]	48 1	26	2127
441		1532	1 768	gi 142863	replication initiation protein [Bacillus subtilis] pir B26580 B26580 replication initiation protein - Bacillus ubtilis	48	23	765
444	5 3	3898	5298	gi 145836		48	24	1401
484	2 1 3	388	1110	gi 146551		48	18	723
542	3 - 1	1425	1 2000	pir S28969 S289	IN-carbamoylsarcosine amidohydrolase (EC 3.5.1.59) - Arthrobacter sp.	48	27	576
566		e .	1019	gi 153490		48	24	1017
611	-	2	730	gi 1103507	unknown [Schizosaccharomyces pombe]	48	38	729
624	 -	1255	1 665	gi 144859	ORF B [Clostridium perfringens]	48	26	591
846	-	1014	1 508	gi 537506	paramyosin [Dirofilaria immitis]	48	27	507
1020		99	950	gi 1499876	magnesium and cobalt transport protein [Methanococcus jannaschii]	48	30	885
1227		-	174	gi 493730	lipoxygenase [Pisum sativum]	48	35	174
1266		·	405	gi 882452 	ORF f211; alternate name yggA; orf5 of X14436 [Escherichia coli] gil41425 ORF5 (AA 1-197) [Escherichia coli] (SUB 15-211)	48	24	405
2071	1 1 7	707	381	gi 1408486	HS74A gene product [Bacillus subtilis]	48	25	327
2398	1 4	463	1 233	gi 1500401	reverse gyrase [Methanococcus jannaschii]	48	40	231
2425	1 4	476	1 246	pir H48563 H485	G1 protein - fowlpox virus (strain HP444) (fragment)	48	40	231
2432	1 4	446	1 225	gi 1353703	Trio [Homo sapiens]	48	33	222
2453	1 7	794	399	gi 142850	division initiation protein [Bacillus subtilis]	48	29	396
2998	1 - 4	469	1 236	gi 577569	PepV [Lactobacillus delbrueckii]	48	31	234

Table

1410 1689 1011 273 1086 945 1065 267 405 192 486 009 432 744 849 942 549 249 831 702 963 480 519 816 | % ident | length (nt) 32 24 26 27 31 24 38 59 34 28 32 19 17 59 28 23 25 28 28 47 47 47 47 47 47 47 47 47 47 47 47 47 47 | & sim 407 teredicted membrane helices, homology with B. subtilis men Orf3 Rowland et. al. unpublished Accession number M74183), approximately 1 minutes on updated Rudd map; putative [Escherichia coli] sp|P37355|YFBB_ECOLI HYPOTHETICAL 26.7 KD PROTEIN IN MEND-MENB ORF 5' of ECRF3 [herpesvirus saimiri HVS, host-squirrel monkey, eptide, |quinolone resistance norA protein protein [Methanococcus jannaschii] Possible operon with orfG. Hydrophilic, no homologue in the atabase; 126.7% of identity in 165 aa to a Thermophilic bacterium hypothetical | protein 6; putative (Bacillus subtilis) pir|S51177|S511 |trans-activator protein - Equine infectious anemia virus myo-inositol transporter [Schizosaccharomyces pombe] |D9509.27p; CAI: 0.12 [Saccharomyces cerevisiae] G-box binding factor [Dictyostelium discoideum] |hypothetical EcsB protein (Bacillus subtilis) C33G8.2 gene product [Caenorhabditis elegans] D-alanine racemase cds [Bacillus subtilis] pps1; B1496_C2_189 [Mycobacterium leprae] processing protease [Bacillus subtilis] |ORF YKL094w (Saccharomyces cerevisiae] |unknown [Mycobacterium tuberculosis] |hippuricase [Haemophilus influenzae] |PC4-1 gene product [Bradysia hygida] [esterase [Acinetobacter lwoffii] | 140 kDa protein [Plasmid pJM1] putative [Bacillus subtilis] [butative [Escherichia coli] |YqkI (Bacillus subtilis] [YqjV [Bacillus subtilis] |mucin [Homo sapiens] I match gene name acession |gi|1303989 |gi|1511555 1gi11204835 gi|1420856 |gi|1255425 |gi|1303973 |gi|1209223 1gi | 1403455 |gi|1256621 |gi|145836 |gi|11177254 |gi|945219 |gi|540083 |gi|438466 |gi|927340 |gi|486143 |gi|516608 |gi|456562 |gi|142824 |gi|150756 |gi|466882 |gi|243353 |gi|142822 6710 1 4279 1676 6925 8863 11174 1093 3107 1 2232 1084 1884 115108 1 2773 1257 1 280 400 831 301 943 483 633 819 Contig | ORF | Start | ID | ID | (nt) | 2145 2321 3641 1 9135 1 2178 1884 1 2620 7524 116118 7141 5022 1 2022 1109 1 2072 1634 1115 14 1 599 196 152 962 1 492 7 -7 -112 - 2 - 2 110 7 -_ _ <u>~</u> 'n 7 _ _ _ --122 3042 3686 4027 24 43 44 69 81 263 529 565 692 168 404 654 391

Table 2

348 753 396 291 219 3990 936 1068 1260 396 561 1209 1209 1716 228 594 282 537 861 615 59 30 35 25 56 35 35 35 30 30 28 28 25 46 46 46 46 46 46 46 46 46 46 l % sim 46 46 |cerebellar-degeneration-related antigen (CDR34) [Homo sapiens] gil182737 orf4; putative transporter; Method: conceptual translation supplied by lenvelope glycoprotein C2V3 region (Human immunodeficiency virus type pir N 29770 N 29770 cerebellar degeneration-related protein - human |single-stranded-DNA-specific exonuclease [Haemophilus influenzae] |dihydrolipoamide acetyltransferase [Haemophilus influenzae] cerebellar degeneration-associated protein [Homo sapiens] |hypothetical protein SP:P45869 [Methanococcus jannaschii] proton antiporter efflux pump [Mycobacterium smegmatis] [Aspartate aminotransferase [Bacillus circulans] [polymorphic antigen [Plasmodium falciparum] (urea?) amidolyase [Haemophilus influenzae] [ATP-dependent DNA ligase [Candida albicans] |myosin heavy chain [Entamoeba histolytica] [ATP-binding protein [Bacillus subtilis] |pir|S51910|S519 |G4 protein - Sauroleishmania tarentolae |clumping factor [Staphylococcus aureus] |portal protein [Bacteriophage SPP1] [Yel040p [Saccharomyces cerevisiae] author [Mycobacterium smegmatis] |F54D5.7 [Caenorhabditis elegans] [putative [Escherichia coli] ORF f286 [Escherichia coli] |ORF_f181 [Escherichia coli] |EF [Streptococcus suis] |pacA [Bacteriophage P1] acession |gi|1511057 |gi|1204449 |gi|1523812 |gi|1041334 1gi11110518 |gi|1147557 |gi|1246901 |gi|1221884 |gi|1197634 |gi|397526 gi | 558073 |gi|755153 |gi|145836 |gi|603639 1gi 1298032 |gi|180189 1gi1607573 |gi|537052 |gi|215635 |gi|537148 |gi|305080 |gi|15470 | 1 | 2419 | 1211 | 4 | 3941 | 7930 1 4093 14 | 4719 | 3652 8284 14 | 1876 | 2490 1 1 1 2663 1 1455 | 2 | 649 | 1242 1285 4361 3058 110300 1 862 1 4192 753 563 1 398 Contig |ORF | Start | ID |ID | (nt) | 1 4 | 5028 1 5 1 2477 1 6 | 4708 14 | 2777 9543 1 3 | 1512 110518 211 1351 | 1 1793 3 1 437 1 1 485 1 - 1 1 1 1 -1-3 11 1 2 7 1 1076 61 170 253 384 132 191

Table

S. aureus - Putative coding regions of novel proteins similar to known proteins

+			+		+ 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			+
Contig	IORF IID	Start (nt)	Stop (nt)	match acession	match gene name	e sin	% ident	length (nt)
3685			402	gi 450688	hsdM gene of EcoprrI gene product [Escherichia coli] pir 538437 538437 hsdM protein - Escherichia coli pir 509629 509629 hypothetical protein A - Escherichia coli SUB 40-520)	46	33	405
4176		673	338	gi 951460	FIM-C.1 gene product [Xenopus laevis]	46	31	336
1 37		1 4813	5922	gi 606064	ORF_f408 [Escherichia coli]	45	24	1110
38	116	111699	112004	gi 452192		45	24	306
1 87	- 5	1748	1 2407	gi 1064813	homologous to sp:PHOR_BACSU [Bacillus subtilis]	45	23	099
103	-112	114182	113385	gi 1001307	hypothetical protein [Synechocystis sp.]	45	22	798
112	114	114791	113811	gi 1204389	H. influenzae predicted coding region HI0131 [Haemophilus influenzae]	45	23	981
145	4	4483	3461	gi 220578	lopen reading frame [Mus musculus]	45	20	1023
170	9	6329	1 4965	gi 238657 	AppC=cytochrome d oxidase, subunit I homolog [Escherichia coli, K12, eptide, 514 aa]	45	27	1365
1 206	- 5	1 5230	1 4346	gi 1222056	aminotransferase [Haemophilus influenzae]	45	27	882
1 228		09	1 716	gi 160299	glutamic acid-rich protein [Plasmodium falciparum] pir A54514 A54514 glutamic acid-rich protein precursor - Plasmodium alciparum	45	23	657
1 288	-	2	1015	gi11255425	[C33G8.2 gene product [Caenorhabditis elegans]	45	23	1014
313	8	1 4339	3128	gi 581140	NADH dehydrogenase [Escherichia coli]	45	30	1212
332	-	914	1 459	gi 870966	F47A4.2 [Caenorhabditis elegans]	45	20	456
344		ю —	1 221	gi 171225	kinesin-related protein [Saccharomyces cerevisiae]	45	26	219
441	2	1501	1073	gi 142863 	replication initiation protein [Bacillus subtilis] pir B26580 B26580 replication initiation protein - Bacillus ubtilis	45	27	429
672		1 2	1 982	gi 1511334	M. jannaschii predicted coding region MJ1323 [Methanococcus jannaschii]	45	22	981
1 763	- 3	1345	851	gi 606180	ORF f310 [Escherichia coli]	45	24	495
886	ю 	379	846	gi 726426 	similar to protein kinases and C. elegans proteins F37C12.8 and 37C12.5 [Caenorhabditis elegans]	45	30	468
948		e 	473	gi 156400 	myosin heavy chain (isozyme unc-54) [Caenorhabditis elegans] pir A93958 MWKW myosin heavy chain B - Caenorhabditis elegans sp P02566 MYSB_CAEEL MYOSIN HEAVY CHAIN B (MHC B).	45	25	471
1158		5	376	gi 441155	ransmission-blocking target antigen [Plasmodium falciparum]	45	35	375
1 2551		4	1 285	gi 1276705	ORF287 gene product [Porphyra purpurea]	45	28	282
1 3967		1 42	374	lgi1976025	HrsA [Escherichia coli]	45	28	333
				, 11 11 11 11 11 11 11 11 11 11				

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

Contig	IORF	Start (nt)	Stop (nt)	match acession	match gene name	e sin	* ident	1ength (nt)
1 52	7	6931	5846	lgi1467378	unknown [Bacillus subtilis]	44	22	1086
1 138	8	6475	6849	gi 173028	thioredoxin II (Saccharomyces cerevisiae]	44	28	375
1 221	5	7032	1 5617	gi 153490	tetracenomycin C resistance and export protein [Streptomyces laucescens]	44	21	1416
1 252	7	1331	1122	gi 1204989	hypothetical protein (GB:U00022_9) [Haemophilus influenzae]	44	30	210
1 263	2	3265	1 2093	gi 1136221	carboxypeptidase [Sulfolobus solfataricus]	44	26	1173
365	4	4963	3524	gi 1296822	orfl gene product [Lactobacillus helveticus]	44	31	1440
1 543	e 	1315	1833	gi 1063250 		44	24	519
544	4	3942	4892	gi 951460	FIM-C.1 gene product [Xenopus laevis]	44	32	951
792		1224	613	gi 205680	high molecular weight neurofilament [Rattus norvegicus]	44	28	612
44	118	11303	111911	gi 1511614 	molybdopterin-guanine dinucleotide biosynthesis protein A [Methanococcus jannaschii]	43	27	609
1 59	8	3665	1 5128	gi 153490	tetracenomycin C resistance and export protein [Streptomyces laucescens]	43	21	1464
59	100	5536	1 7527	gi 153022		43	22	1992
66	-	1346	1 681	gi 1419051	unknown [Mycobacterium tuberculosis]	43	21	999
310	80	9402	112134	gi 397526	clumping factor [Staphylococcus aureus]	43	21	2733
1 432		2782	1 2303	pir A60540 A605	Isporozoite surface protein 2 - Plasmodium yoelii (fragment)	43	29	480
519	m ——	2547	3122	sp Q06530 DHSU_	ISULFIDE DEHYDROGENASE (FLAVOCYTOCHROME C) FLAVOPROTEIN CHAIN PRECURSOR (EC 1.8.2) (EC) (FCSD).	43	23	576
4	113	12053	13321	gi 295671	selected as a weak suppressor of a mutant of the subunit AC40 of DNA ependant RNA polymerase I and III (Saccharomyces cerevisiae)	42	18	1269
94	1 2 1	1768	1091	lgi1501027	ORF2 [Trypanosoma brucei]	42	31 +	678
127	4	1 5791	1 4550	gi 42029	ORF1 gene product [Escherichia coli]	42	21	1242
1 297	3	1515	1036	gi 142790	(ORF1; putative (Bacillus firmus)	42	25	480
344	9	4097	3525	gi 40320	ORF 2 (AA 1-203) [Bacillus thuringiensis]	42	30	573
1 512	-	1 2167	1115	gi 405957	yeeF [Escherichia coli]	42	23	1053
631		2434	1223	gi 580920 	rodD (gtaA) polypeptide (AA 1-673) [Bacillus subtilis] pir S06048 S06048	42	24	1212

Table 2

S. aureus - Putative coding regions of novel proteins similar to known proteins

+			÷					+ -
Contig	ID	Start (nt)	Stop (nt)	match acession	match gene name	E S S	* 1dent	length (nt)
1 685	. 3	1 2359	1 1739	gi 1303784	YqeD [Bacillus subtilis]	42	19	621
4132		1 787	395	lgi11022910	protein tyrosine phosphatase (Dictyostelium discoideum)	42	25	393
86	5	1375	884	gi 309506	spermidine/spermine N1-acetyltransferase [Mus saxicola] pir S43430 S43430 spermidine/spermine N1-acetyltransferase - spiny ouse (Mus saxicola)	41	30	492
191	112	114797	114075	gi 1124957	orf4 gene product [Methanosarcina barkeri]	41	22	723
1 212	9 -	1 2150	1 3127	gi 15873	observed 35.2Kd protein [Mycobacteriophage 15]	41	26	978
213	3	1263	1 2000	gi 633692		41	18	738
408	4	1 2625	3386	gi 1197634 	orf4; putative transporter; Method: conceptual translation supplied by author [Mycobacterium smegmatis]	41	24	762
1 542	1 - 1	. 3	1103	gi 457146	rhoptry protein [Plasmodium yoelii]	41	21	1101
1 924	-	7	1 475	pir JH0148 JH01	.rlJH0148jJH01 nucleolin - rat	41	30	474
1562			1 402	gi 552184 	asparagine-rich antigen Pfa35-2 [Plasmodium falciparum] pir \$27826 \$27826 asparagine-rich antigen Pfa35-2 - Plasmodium alciparum (fragment)	40	20	402
1 2395	1 1	518	261	pir S42251 S422	hypothetical protein 5 - fowlpox virus	40	18	258
4077		m 	1 305	gi 1055055 	lcoded for by C. elegans cDNA yk37g1.5; coded for by C. elegans cDNA yk5c9.5; coded for by C. elegans cDNA yk1a9.5; alternatively spliced form of F52C9.8b [Caenorhabditis elegans]	39	21	303
928		1003	503	gi 1255425	[Canonitis elegans]	37	25	501
65	112	1 8294	110636	gi 535260		36	24	2343
63	5	1 3550	6208 1	gi 298032	EF [Streptococcus suis]	36	19	4530
1 544	3	1 2507	1 3601	gi 1015903	ORF YJR151c [Saccharomyces cerevisiae]	35	22	1095
63	4	1949	3574	gi 552195 		32	27	1626